

This document contains treatment criteria for use of:

Page 003: Section A: Cancer drugs/indications currently funded by the Cancer Drugs Fund (CDF)

Page 042: Section B: NICE & NHSE approved cancer drugs/indications routinely funded by NHSE from 1st April 2016

Page 280: Section C: NHS England interim cancer treatment options funded during the COVID-19 pandemic

ver1.379

26-Nov-25

Opera	tions and Informatio	n Spe	cialised Commissioning					
	& Corp. Ops.		nmissioning Strategy					
			3 3,					
s Gateway Reference		056	05					
Purpose Policy								
Name Nation	al Cancer Drug Fur	d List						
NHS E	ngland Cancer Dru	gs Fund Tea	am					
Date 29 July								
Target Audience Foundation Trust CEs., Medical Directors, NHS England Regions Directors, NHS England Directors of Commissioning Operation Directors of Finance, NHS Trust CEs, Patients; Patient Group: Charities; Pharmaceu								
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#### A. National CDF List

This list should be read in conjunction with 'Appraisal and Funding of Cancer Drugs from July 2016 (including the new Cancer Drugs Fund) A new deal for patients, taxpayers and industry' published by NHS England on 8 July 2016 at www.england.nhs.uk/ourwork/cancer/cdf

				Availab	ble to new	patients		Transition	Eligible for	Interim Funding agreed by	CDF Managed	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Interim Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
AVE3	Avelumab in combination with axitinib	For use in treatment-naïve patients with advanced renal cell carcinoma where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with the combination of avalumab and autinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic autinometr therapy.  2. The prescribing clinician is fully aware of the management of and the breatment modifications that may be required for immune-related adverse reactions due to checkpoint inhibitor treatments including presumonits, colitis, reporting, endocrinospathins, hepatitis and other immune-related adverse reactions.  3. The patient has unresectable locally advanced or metastatic renal cell carcinoma (RCC) which has either a clear cell component or is one of the types of RCC as indicated below. Please indicate below which RCC historiogy applies to this patient:  8. RCC with a clear cell component or Papilary RCC or Companyblo RICC or Companyblo RI		rom 31-Jul-2(	20	No	n/a	Yes	Agreed	Yes	nca

				Availa	ble to new	patients		Transition	Eligible for	Interim Funding agreed by	CDF Managed	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Interim Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
AXI02a_v1.0	Axicabtagene ciloleucel	chemoimmunotherapy ARD who would otherwise be intended for potential stem cell transplantation where the following criteria are met:  Insi form is for the approval of Jeucapheresis and manufacture of CAR-T cells. There is a second part to this form which relates to the subsequent injusion of CAR-T cells and this will be available dries under the completed day of the form (AVOZD) can only be completed as a continuation of this first part. The other form (AVOZD) can only be completed as a continuation of this first part. The form	1. This papellation is being mode by and that locaphrenis for and treatment with auxiliarged collected modified CART cells will be initiated by a consultant hematodispit or medical accordings to predict the relation of the treating Proof. SEAC. And MCRCC. CART cell and disciplinary team.  The patient is auxiliary like gift years or own of the date of garwing the auxiliary and according to the treatment of the treatment with a scientific or the treatment of the treatment of the treatment with a scientific or the treatment of the treatment with a scientific or the treatment of the treatment with a scientific or the treatment of the treatment with a scientific or the treatment of the treatment with a scientific or the treatment of the tre		From 27-Apr-	223	No	n/a	Yes	Agreed	Yes	NCA

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				Availab	ole to new	patients				Interim Funding	CDF	
							Transition	Transition Funding agreed	Eligible for	agreed by	Managed	Expected Entry
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Drug (Old CDF) Indication (Yes or No)	by manufacturer (Agreed, Rejected, Pending)	Funding (Yes, No, Not currently applicable (NCA))	(Agreed, Rejected, Pending, Not currently applicable (NCA))	Scheme (Yes, No, Not currently applicable (NCA))	into Baseline Commissioning (Date if known or Not currently applicable (NCA))
AXI02a_v1.0	Axicabtagene ciloleucel	in patients who relapse within 12 months of completion of 1st line chemoimmundherapy AND who would otherwise be intended for potential stem cell transplantation grow has refractory to 1st line chemoimmunotherapy AND who would otherwise be intended for potential stem cell transplantation where the following criteria are met:  This farm is for the approval of leucaphnesis and manufacture of GAH 7 cells. There is a second part to this form which relates to the subsequent following criteria are related to the control of the complete of the control of the control of the control of the farm (AND 2d) can only be completed as a continuation of this first part of the farm (AND 2d) can only be completed as a continuation of this first part of the farm (AND 2d) and must be completed on influsion of CAH 7 cells.	13. The patient has an ECOG performance score of 0 or 1. Please enter below as to the patient's current ECOG performance status (PS): The ECOG performance status scale is as follows: 15 of the patient is fully active and able to carry oal pre-disease performance without restriction 15 of the patient is restricted in physically streuous activity but is ambidatory and able to carry oat work of a light or sedentary nature eg light housework, office work 15 of the patient is restricted in physically streuous activity but is ambidatory and able to carry out any work activities and is up and about more than 50% of waking hours 15 of the patient is completely disabled, cannot carry out any selfcare and is totally confined to be do rot waiting hours 15 of the patient is completely disabled, cannot carry out any selfcare and is totally confined to be do rot hair 15 patient is completely disabled, cannot carry out any selfcare and is totally confined to be do rot hair 16 patient currently has a performance status of either 16 of 90 or 16 of 90 or 16 of 90 or 16 of 90 or 17 of 90 or 18 of 90 or 18 of 90 or 19 of 90	F	From 27-Apr-	23	No	n/a	Yes	Agreed	Yes	NCA
AXIO2b_v1.0	Axicabtagene ciloleucel	Axicabtagene ciloleucel for treating relapsed/refractory diffuse large B-cell lymphoma (DLBCL) or high-grade B-cell lymphoma (ol.BCL) or high-grade B-cell lymphoma and in adult patients either, who relapse within 12 months of completion of 1st line chemoimmunotherapy AND who would otherwise be intended for potential stem cell transplantation gw hoa re efractory to 1st line chemoimmunotherapy AND who would otherwise be intended for potential stem cell transplantation where the following criteria are met:  This second part of the form is to document the date of infusion of CAR-T cell therapy and for registration of this infusion with NHS England as that the treating Trust is reimbursed for the cost of axicabtagene ciloleucel. There is a first part of the form for the approval of leucaphresis and manufacture of CAR-T cells which has already been completed (AXIO2a). This second part of the form (AXIO2b) should only be completed as a continuation form once the date of CAR-T cell infusion is known.	1. This application for continuation is being made by and treatment with axicabtagene ciloleucel-modified CAR-T cells will be initiated by a consultant haematologist/medical oncologist specifically trained and accredited in the use of systemic anti-cancer therapy and working in an accredited CAR-T cell treatment centre and who is a member of the National CAR-T clinical Panel for DLSCL and H6BCL and AR-T cell multideoliphany teams.  2. The patient has an ECOG performance score of 0 or 1 or 2. Please tick one of the box below as to the patient's current ECOG performance status (PS):  The ECOG performance status scale is a so follows:  PS 0 The patient is a so follows:  PS 1 The patient is a so follows:  PS 1 The patient is a so follows:  PS 1 The patient is stricticed in physically strenuous activity but is ambilatory and able to carry out any work activities and is up and about more than 50% of waking hours  PS 2 The patient is capable of only limited selferar els confined to bed or chair more than 50% of waking hours  PS 4 The patient is capable of only limited selferar els confined to bed or chair more than 50% of waking hours  PS 4 The patient is capable of only limited selferar els confined to bed or chair The patient currently has a performance status of:  -ECOG PS 1 or  -ECOG PS 2 or  -ECOG PS 2 or  -ECOG PS 2 or  -ECOG PS 2 or  -ECOG PS 3 or  -ECOG PS 4 or  -ECOG PS 5 or  -ECOG PS 5 or		from 27-Apr-	23	No	n/a	Yes	Agreed	Yes	NCA

				Availab	le to new p	patients				Interim Funding	CDF	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Transition Funding agreed by manufacturer (Agreed, Rejected, Pending)	Eligible for Interim Funding (Yes, No, Not currently applicable (NCA))	agreed by manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Managed Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
BELA1	Belantamab mafadotin in combination with bortezomib and dexamethasone	Belantamab mafadotin in combination with bortezomib and dexamethasone as 2nd line treatment of relapsed or refractory myeloma in adult patients who previously receives lenal diomide as part of 1st line systemic therapy where the following criteria have been met:	In this application for betalantamb transferdin in combination with borteomic and decamethacone is being made by and the first cycle of systemic anti-cancer therapy with belantamab will be prescribed by a combinating specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patients with amyliodois or POEMS syndrome are not eligible for belantamab marfactors.  Note: patients with amyliodois or POEMS syndrome are not eligible for belantamab marfactors.  Note: patients with amyliodois or POEMS syndrome are not eligible for belantamab marfactors.  Note: patients with amyliodois or POEMS syndrome are not eligible for belantamab marfactors.  Note: patients with amyliodois or POEMS syndrome are not eligible for belantamab marfactors.  Note: patients with amyliodois or POEMS syndrome are not eligible for belantamab marfactors.  Note: patients with amyliodois or POEMS syndrome are not eligible for belantamab marfactors.  Note: patients with a syndrome and patients with a patient syndrome and patients which syndrome and patients with a patient syndrome and patients which syndrome and patients with syndrome and patients which syndrome and patients with a patient syndrome and patients who have been done or patients who have been previously treated with a 1st line lenalidomide-containing regimen which is commissioned by NHS England or is part of a 1st line treatment regime in a NHH-badded clinical trial.  Please confirm below which 1st line treatment was received by the patients.  4. This patient has been previously treated with a 1st line lenalidomide-containing regimen which is commissioned by NHS England or is part of a 1st line treatment regimen in a NHH-badded clinical trial.  Please confirm below which 1st line treatment was received by the patients.  4. This patient has been previously treated with a 1st line benefit of stronger and eligible disease		rom 12-Jun-2	15	No	nca	Yes	Agreed	No	nca

Blueteq Form ref:				Availab	ole to new p	atients		Transition	Eligible for	Interim Funding agreed by	CDF Managed	
	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Interim Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
BELA1	Belantamab mafadotin in combination with bortezomib and dexamethasone	Belantamab mafadotin in combination with bortezomib and dexamethasone as 2nd line treatment of relapsed or refractory myeloma in adult patients who previously received lenalidomide as part of 1st line systemic therapy where the following criteria have been met:	11. Belantamab mafodotin will be used only in combination with bortezomib and dexamethasone and not with any other anti-myeloma agents.  12. The prescribing clinician is aware of the risk of corneal adverse reactions with belantamab mafodotin and that an ophthalmic examination including visual acuity and slit lamp examination must be performed by an eye care professional prior to each of cycles 1, 2, 3 and 4 and then during treatment as indicated.  13. Arrangements have been put in place for the eye care professional to categorize both the degree of any corneal damage and the best corrected visual acuity in the most severely affected eye and for these results to be communicated to the myeloma team.  14. Since belantamab mafodotin dose modifications are partly based on corneal examination findings and/or changes in best corrected visual acuity, the patient's ophthalmic examination findings will be reviewed before dosing and will determine the belantamab mafodotin dose based on the highest category from the corneal examination and/or best corrected visual acuity finding in the most severely affected eye.  15. The patient will be advised to administer preservative-free artificial tears for use at least 4 times daily throughout the time of treatment with belantamab mafodotin.  16. The patient will be treated with belantamab mafodotin until disease progression or the occurrence of excessive toxicity or the withdrawal of patient consent, whichever is the sooner.  18. A formal medical review as to how belantamab mafodotin is being tolerated and whether treatment with belantamab should continue or not will be scheduled to occur after each of the first 4 cycles of treatment.  19. The prescribing clinician understands that given the potentially necessary frequency and duration of treatment breaks during treatment with belantamab mafodotin, this indication is exempt from NHS England's treatment break policy.  Note: if there is disease progression during a treatment break from belantamab mafodotin, the during's Summary of	f	From 12-Jun-25	5	No	nca	Yes	Agreed	No	nca

				Availal	ble to new	patients				Interim Funding	CDF	
				7.7.5	J.C 10 1.C11	putients		Transition	Eligible for	agreed by	Managed	Expected Entry
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)		Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Interim Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	into Baseline Commissioning (Date if known or Not currently applicable (NCA))
BELZUT1a	Belzutifan monotherapy	For adult patients with von Hippel-Lindau (VHL) disease who require systemic therapy for VHL associated renal cell carcinoma, central nervous system haemangioblastomas or pancreatic neuroendocrine tumours, AND for whom localised procedures are unsuitable or undesirable where the following criteria have been met:  This form BELZUTIa is for the FIRST ever application for a patient to commence belzutfian for the above indication. The form BELZUTIb is for either continuation of bezutfan beyond disease progression in one dominant tumour but with continued benefit in other equally dominant VHL associated tumour to the one which previously resulted in the original indication for belzutifian for a different VHL associated tumour to the one which previously resulted in the original indication for belzutifian for a different VHL associated tumour to the one which previously resulted in the original indication for belzutifian for a different VHL associated tumour to the one which previously resulted in the original indication for belzutifian for a subsequent restart of procedures are unsultable or undesirable.	1. This application is both being made by and the first cycle of systemic articisance therapy. 2. The patient is an adult with a VIII, germline alteration.  Please state the type of VIII. decimine alteration including on system therapy with beating or whether the patient has vivil type 2 disease.  1. The patient's case has been discussed at a VIII mutilissicalizary team meeting which has recommended the use of belautifien for a VIII. associated renal cell carcinoma or a CMS hammaglobatoms or a parcental enumeration including on system therapy with beating or whether the patient has mutilitystem involvement which its the obiniman includination for yetamic therapy with beating or whether the patient has mutilitystem involvement which requires systemic therapy for more than one type of VIII. cancer and for which 2 or more unsuitable or undesirable.  1. The obiniman includination for treatment with belatifier is for CMS hammaglobatoms with or without other VIII. associated tumours which are not yet indicated for localised treatment or the obiniman includination for treatment with belatifier is for CMS hammaglobatoms with or without other VIII. associated tumours which are not yet indicated for localised treatment with patient which the patient was also an advantage of the patient and including the patient was also an advantage of the patient and clinicate to be usualized for localised treatment with patient was also an advantage of the patient and clinicate to be usualized for localised treatment with patient was also an advantage of the patient and clinicate to be usualized or undesirable.  1. In the absenc		From 05-Sep-	24	No	nca	Yes	Agreed	Yes	nca

				Availal	ble to new	patients		Transition	Eligible for	Interim Funding agreed by	CDF Managed	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Interim Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
BELZUTIa	Belzutifan monotherapy	For adult patients with von Hippel-Lindau (VHL) disease who require systemic therapy for VHL asociated renal cell carriomas, central nervous system haemagioblastomas or pancreatic neuroendocrine tumours. AND for whom localised procedures are unsuitable or undesirable where the following criteria have been met:  This form BELZUTIa is for the FIRST ever application for a patient to commence betutifan for the above indication. The form BELZUTIb is for either continuation of betutifan beyond disease progression in one dominant tumour but with continued benefit in other equally dominant VHL associated tumours or a subsequent restart of betutifan for a different VHL associated tumour to the one which previously resulted in the original indication for belzutifan treatment, and for which localised procedures are unsuitable or undesirable.	with a localised procedure for this progressing tumour in there has nevertneess been continued behalf and outside procedure for the progressing tumour in the above the head for an unsatiable/undestrable localised procedure. In such a patient, blueted for state, Italy the behalf and the description of the behalf and the state of the sta		From 05-Sep-	24	No	nca	Yes	Agreed	Yes	nca

				Availa	ible to ne	w patient	s	Transition	Eligible for	Interim Funding agreed by	CDF Managed	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (bu notice or remove served	of al No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
BELZUTIb	Beizutifan monotherapy	For adult patients with von Hippel-Lindau (VHL) disease who require ETHER continuation of belzutfan beyond disease progression in one dominant tumour but who have continued benefit in other equally dominant VHL associated tumour of the noe which previously resulted in the original indication robe learning the teatment, and AND for which localised procedures are unsuitable or undesirable where the following criteria have been met:  The Form BELZUTIa is for the FIRST ever application for a patient to commence betutuffan for a VHL associated tumour of which localised procedures are unsuitable or undesirable. This BELZUTI for its for either continuation of belzutfan beyond disease progression in one dominant tumour but with continued benefit in other equally dominant VHL associated tumour to the one which previously resulted in the indication for belzutfan treatment, and for which localised procedures are unsuitable or undesirable.	- The dominant indication for treatment with belzutifan is for pancreatic neuroendocrine tumour with or without other VHL associated tumours which are not yet indicated for localised treatment  - this patient has multisystem disease with 2 or more of these 3 VHL associated types of cancer which are currently equally dominant as to the need for localised treatment procedures  5. In the absence of systemic therapy with belzutifan the patient would otherwise proceed to treatment for VHL associated tumour(s) with a localised procedure/procedures which is/are considered by the patient and clinician to be unsuitable or undesirable.  Please tick the box below as to the type of localised treatment which would otherwise be employed (surgery or abilative procedure or radiotherapy) and then state the procedure(s) in		From 05-Se	p-24	No	nca	Yes	Agreed	Yes	nca

				Availat	ole to new p	atients		Transition	Eligible for	Interim Funding agreed by	CDF Managed	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Interim Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
BELZUTIb	Belzutifan monotherapy	For adult patients with von Hippel-Lindau (VHL) disease who require ETIHER continuation of belautifan beyond disease progression in one dominant tumour but who have continued benefit in other equally dominant VHL associated tumour to a different VHL associated tumour to the one which localised procedures are unsuitable or undesirable where the following criteria have been met:  The Form BELZUTIa is for the FIRST ever application for a patient to commence behautifan for a VHL associated tumour for which localised procedures are unsuitable or undesirable. This BELZUTIa is for the FIRST ever application for a patient to commence behautifan for a VHL associated tumour for which localised procedures are unsuitable or undesirable. This BELZUTIa form is for either continuation of belzutifan beyond disease progression in one dominant tumour but with continued benefit in other equally dominant VHL associated tumour to the one which prevolusy resulted in the indication for belzutifan for a different VHL associated tumour to the conditional continuation of the condition of the procedures are unsuitable or undesirable.	- performance status 1 or - performance status 2  12. Belzutifan is only to be used as monotherapy for treating VHL associated RCC and/or CNS haemangioblastoma and/or pNET.  13. For the dominant indication/tumour belzutifan is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment or the occurrence of an intervention with a localised procedure for that dominant indication/tumour. Note: belzutifan cannot be restarted for patients who suffer unacceptable toxicity or choose to stop treatment. Patients in such circumstances should be counselled that belzutifan cannot be restarted.  Note: the intention to treat with belzutifan must be with a planned and continued administration of belzutifan until disease progression or unacceptable toxicity or patient choice to stop treatment or the occurrence of an intervention with a localised procedure. Belzutifan is not funded to be used electively in an intermittent treatment schedule with planned "treatment bolidass".		From 05-Sep-2	4	No	nca	Yes	Agreed	Yes	nca

				Avai	lable to ne	v patients			-11 11 C	Interim Funding	CDF	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (bi notice remov served	of No	Transition Drug (Old CDF) Indication (Yes or No)	Transition Funding agreed by manufacturer (Agreed, Rejected, Pending)	Eligible for Interim Funding (Yes, No, Not currently applicable (NCA))	agreed by manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Managed Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
KTE01a_v1.2	Brexucabtagene autoleucel (formerly known as KTE-X19	For treating mantle cell lymphoma (MCL) in adults previously treated with two or more lines of systemic therapy where the following criteria have been met:  This form is for the approval of leucapheresis and manufacture of CAR-T cells. There is a second part to this form which relates to the subsequent infusion of CAR-T cells and this will be available after submission of the first part. The second part of the form (KTEOLB) and must be completed on infusion of CAR-T cells otherwise the treating Trust will not be reimbursed for the cost of brexucabtagene autoleucel.	- nas nad autorogous Sc.T. or - has had allogeneic SCT  8. The patient has been previously treated for MCL with a Bruton's tyrosine kinase (BTK) inhibitor (such as ibrutinib or acalabrutinib) and that the patient progressed either during treatment or following discontinuation of the BTK inhibitor.		From 19-Ja	h-21	No	nca	Yes	Agreed	Yes	nca

				Availab	ole to new p	patients		Transition	Eligible for	Interim Funding agreed by	CDF Managed	
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			1. This application for continuation is made by and treatment with brexucabtagene autoleucel (formerly known as KTE-X19-modified CAR-T) will be initiated by a consultant haematologist/medical oncologists specifically trained and accredited in the use of systemic anti-cancer therapy and working in an accredited CAR-T cell treatment centre and who is a member of the National CAR-T Cilinical Panel for MCL and a member of the treating Trust's MCL and CAR-T cell multidisciplinary teams.  2. The patient has an ECOG performance score of 0 or 1 or 2. Please tick one of the boxes below as to the patient's current ECOG performance status (PS):  The ECOG performance status scale is as follows:									
		For treating relapsed/refractory mantle cell lymphoma (MCL) in patients aged 18 years and over where the following criteria have been met:  This second part of the form is to document	PS 0 - The patient is fully active and able to carry on all pre-disease performance without restriction PS 1 - The patient is restricted in physically strenuous activity but is ambulatory and able to carry out work of a light or sedentary nature e.g. light housework, office work PS 2 - The patient is ambulatory and capable of all selfcare but unable to carry out any work activities and is up and about more than 50% of waking hours PS 3 - The patient is capable of only limited selfcare and is confined to bed or chair more than 50% of waking hours PS 4 - The patient is completely disabled, cannot carry out any selfcare and is totally confined to bed or chair The patient currently has an ECOG performance status of: - ECOG PS 0 or - ECOG PS 1 or - ECOG PS 2									
KTE01b_v1.3	Brexucabtagene autoleucel (formerly known as KTE-X19 (Tecartus®))		3. The patient has either required bridging therapy in between leucapheresis and CAR-T cell infusion or not. Please indicate what type(s) of bridging therapy have been required by ticking the most appropriate option below: - no bridging therapy at all or - corticosteroids only or - ibrutinib monotherapy (only for those patients who previously discontinued a Bruton's tyrosine kinase (BTK) inhibitor without disease progression) or another BTK inhibitor or - radiotherapy only or - radiotherapy only or - corticosteroids and ibrutinib (only for those patients who previously discontinued a BTK inhibitor without disease progression) or corticosteroids and another BTK inhibitor or - corticosteroids and ibrutinib (only for those patients who previously discontinued a BTK inhibitor without disease progression) or corticosteroids and another BTK inhibitor or - corticosteroids and chemo(immuno)therapy or - corticosteroids and chemo(immuno)therapy or - chemo(immuno)therapy and radiotherapy ± corticosteroids	F	From 19-Jan-2	1	No	nca	Yes	Agreed	Yes	nca
			4. The patient does not have known active CNS involvement by the lymphoma.  5. The patient has sufficient end organ function to tolerate treatment with CAR-T cell therapy.  6. Prior to infusion of brexucabtagene autoleucel, 2 doses of tocilizumab are available for use in this patient in the event of the development of cytokine release syndrome.									
			7. Brexucabtagene autoleucel will otherwise be used as set out in its Summary of Product Characteristics (SPC). 8. Following national approval for use of brexucabtagene autoleucel there has been local CAR-T cell multidisciplinary team agreement that this patient continues to have the necessary fitness for infusion and fulfils all the treatment criteria listed here.									

				Availabl	le to new p	patients		Transition	Eligible for	Interim Funding agreed by	CDF Managed	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Interim Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
BREXO1a	Brexucabtagene autoleucel	the following criteria are met:  This form is for the approval of leading the first of the approval of leading the first of CAR-T cells. There is a second part to this form which relates to the subsequent infusion of CAR-T cells and this will be available after submission of the first part. The second part of the form (BREXOIb) can only be completed as a continuation of this first part of the form (BREXOIc) and BREXOID must be completed on infusion	- relapsed disease and ineligible for allogeneic SCT due to comorbid disease (but still fit enough for CAR-T cell therapy with brexucabtagene autoleucel) or contraindicated to allogeneic SCT conditioning or lack of a suitable donor  4. Having fulfilled, and ticked one of the criteria in box 3 above, the patient at the time of demonstration of such refractory/relapsed disease and thus consideration for potential treatment with brexucabtagene autoleucel.  5. The patient does not have an isolated est amendulary ALT elapse is. Life The patient has some autoleucel.  6. At the time of this application for treatment with brexucabtagene autoleucel the patient has soft and a sense of the patient has soft part and the patient has soft part and the patient has soft part and the patient for the patient flows the patient fl		rom 27-Apr-2	23	No	n/a	Yes	Agreed	Yes	NCA
BREXO1b_v1.0	Brexucabtagene autoleucel	Brexucabtagene autoleucel for treating relapsed/refractory Philadelphia negative and positive B cell acute lymphoblastic leukaemia in patients aged 26 years and over where the following criteria are met:  This second form is to document the date of infusion of CAR T cell therapy and for registration of this infusion with NHS England so that the treating Trust is reimbursed for the cost of brexucabtagene autoleucel. There is a first form for the approval of leucapheresis and manufacture of CAR T cells. This second form must use the same unique Blueteq identifier number generated when this patient was registered for leucapheresis and CAR T cell manufacture using the first form	option below: - no bridging therapy at all or - corticosteroids only or - Trill therapy with or without steroids or - straining and record or without steroids or - straining and record or home therapy with or without steroids or	Fre	rom 27-Apr-2	23	No	n/a	Yes	Agreed	Yes	NCA

				Availa	able to new	patients		Torrestation	Flicthia for	Interim Funding	CDF	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice o remova served)	, No	Transition Drug (Old CDF) Indication (Yes or No)	Transition Funding agreed by manufacturer (Agreed, Rejected, Pending)	Eligible for Interim Funding (Yes, No, Not currently applicable (NCA))	agreed by manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Managed Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
DARO3	Darolutamide in combination with androgen deprivation therapy (ADT)	For the treatment of patients with newly diagnosed metastatic hormone-sensitive prostate cancer who are unsuitable for treatment with docetave himser the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with disordurande will be prescribed by a consultant specialist specifically trained and according in the use of systemic anti-cancer therapy.  2. The patient either has a proven histological or cytological diagnosis of adenocarrisons of the prostate or has presented with a clinical picture consistent with metastatic prostate cancer with both widespread bone metastace according the patient of the patient is newly diagnosed metastatic prostate cancer that is hormone sensitive and has currently received androgen deprivation therapy (ADT) for no longer than 3 months before starting an androgen receptor targeted agent.  Please enter below as to which scenario applies to this patient  - the patient has not yet received any ADT for metastatic prostate cancer or the patient has not yet received any ADT for metastatic prostate cancer or the patient has not received any ADT for metastatic prostate cancer or the patient has not received any ADT for metastatic prostate cancer or the patient has not received any ADT for metastatic prostate cancer or the patient has not received any ADT for metastatic prostate cancer or the patient has not received any ADT for metastatic prostate cancer or the patient has not received any ADT for metastatic prostate cancer or the patient has not received any ADT for metastatic prostate cancer or the patient has not received any ADT for metastatic prostate cancer or the patient has not received any ADT for metastatic prostate cancer or the patient has not received any ADT for metastatic prostate cancer or the patient has not received any ADT for metastatic prostate cancer or the patient has not received any ADT for metastatic prostate cancer or the patient has not received any ADT for metastatic prostate cancer or the patient has not received any ADT for metastatic prostate cancer or the patient has not received by ADT for patient has not received any ADT for metastatic prostate cancer or the patient has		From 24-Oct		No	n/a	Yes	Agreed	No	12-Dec-25

				Availa	able to new	patients		Transition	Eligible for	Interim Funding	CDF	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Interim Funding (Yes, No, Not currently applicable (NCA))	agreed by manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Managed Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
DOS1	Dostarlimab	Dostariimab monotherapy for patients with microsatellite instability high (MS-H) or mismatch repair deficient (dMMR) recurrent/davanced endometrial carcinoma after prior platinum-based chemotherapy where the following criteria have been met:	1. This application is being made by and also that the first cycle of systemic anti-cancer therapy with dostarlimab will be prescribed by a consultant specialist specifically trained and accordance in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully wave of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-1/PD-11 treatments including pneumonitis, colitis, nephritis, endocrinopathies, bepatitis and skin toxicity.  3. The patient has prome histological diagnosis of endometrical carcinoma.  Please mark below whether the histology in this patient is endometriol or not:  - the histology is of non-endometriold type  - the histology is of non-endometriold type  - the histology is of non-endometriold type  - the patient previously had a hysterectomy and relapsed with local recurrence only or  - the patient previously had a hysterectomy and relapsed with local recurrence only or  - the patient previously had in hysterectomy and relapsed with both local recurrence and distant disease or  - the patient previously had collar advanced disease, clin of his we surgery and has relapsed with local recurrence and distant disease or  - the patient previously had collar advanced disease, clin of his we surgery and has relapsed with both call recurrence and distant disease or  - the patient previously had locally advanced disease, clin of his we surgery and has relapsed with both call recurrence and distant disease or  - the patient previously had locally advanced disease, clin of his we surgery and has relapsed with the local recurrence and distant disease or the patient surgeriously had locally advanced disease, clin for the surgery and has relapsed with the history of the patient has progressive disease during or following previous platinum-based therapy for recurrent/locally advanced/metastatic endometrial carcinoma.  Place and the patient has no symptomatic brain or leptomeningeal metastases.  - The patient has an ECOS perf		From 08-Feb-	222	No	n/a	Yes	Agreed	Yes	nca

				Availa	able to new	patients				Interim Funding	CDF	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice or removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Transition Funding agreed by manufacturer (Agreed, Rejected, Pending)	Eligible for Interim Funding (Yes, No, Not currently applicable (NCA))	agreed by manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Managed Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
DOS3		For the 1st line treatment of mismatch repair proficient (pMMR) or microsatellite stable endometrial carcinoma in adult patients who have recurrent or primary advanced disease and who are not candidates for potentially curative surgery or radiotherapy but are eligible for systemic therapy where the following criteria have been met:	6. Dostarlimab will be given in combination with carboplatin and paclitaxel unless there is a clear contraindication to the use of one or both cytotoxic agents.  Please mark below which scenario applies to this patient:  - the intent is to use the combination of carboplatin and paclitaxel as the chemotherapy partner to dostarlimab or - the partiest bas a clear contraindication to the use of carboplatin and/or paclitaxel and hence an alternative platinum-based combination therapy must be used as the chemotherapy		From 25-Nov	25	No	n/a	Yes	Agreed	No	tbc

				Availabl	le to new p	atients				Interim Funding	CDF	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Transition Funding agreed by manufacturer (Agreed, Rejected, Pending)	Eligible for Interim Funding (Yes, No, Not currently applicable (NCA))	agreed by manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Managed Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
DURS		instability-high endometrial carcinoma in adult patients who have recurrent or primary advanced disease and who are not candidates for potentially curative surgery or radiotherapy or chemoradiotherapy but are eligible for systemic therapy where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with dural authority of constructions of patients and pacificate will be prescribed by a consciousnia specialist specialist pacification and accordiated in the such optimized of systemic anti-cancer through.  2. The prescribing clinican is fully aware of the management of, and the treatment modifications that may be required for, immune-related adverse reactions due to anti-PD-1 treatments including personnolists, collins, application, propriets, exportion, importation and this toxicity.  3. The patient sha in histologically—or cyclologically confirmed diagnosis of endometrial carcinoma (including clear cell and aerous histological)—or confirmed diagnosis of endometrial carcinoma (including clear cell and aerous histological)—or common of the surface of the patients with carcinosarroma (Niveed Mullerian tumour) are eligible but otherwise uterine sarcomas of any luid are NOT eligible for duraulumab in this indication.  4. The patient's tumour has a documented presence of mismatch repair deficiency (GMMRI) or microsatellistic instability (NSHI) confirmed by validated testing.  5. The patient either has a 1st recurrence of endometrial carcinoma after surgery or radiotherapy or has presented with primary locally advanced or metastatic endometrial carcinoma and in whichever carcinol is not a candidate for any potentially currient treatment with surgery or radiotherapy or chemoradotherapy or the moradotherapy or the patient with primary stage. In the disease and has received no systemic therapy or presented with primary stage life disease and has received no systemic therapy or presented with primary stage life disease and has received no systemic therapy or presented with primary stage life disease and has received no systemic therapy or presented with primary stage life disease and has received no systemic therapy or presented with primary stage life disease and has received no systemic therapy or presented with primary stage l		om 26-Mar-2	5	No	n/a	Yes	Agreed	No	nca

				Availal	ble to new p	patients		Transition	Eligible for	Interim Funding agreed by	CDF Managed	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed,		manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with durvalumab monotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.									
			2. The patient has a histologically or cytologically determined diagnosis of small cell lung cancer (SCLC).									
			3. The patient has limited stage SCLC.									
			4. The patient has been treated with platinum-based chemoradiotherapy (etoposide plus either cisplatin or carboplatin) and there has been no evidence of disease progression following this.									
			Please mark below whether the radiotherapy was concurrent with chemotherapy or sequential after chemotherapy:									
			- concurrent radiotherapy and chemotherapy or									
			- sequential radiotherapy after chemotherapy									
		Durvalumab monotherapy for patients with limited- stage small cell lung cancer	Note: NHS England expects concurrent chemoradiotherapy to be the preferred way of giving platinum-based chemotherapy and radiotherapy in line with the 2019 NICE Clinical Guideline for SCLC.									
DUR7	Durvalumab	whose disease has not progressed	S. The patient has been treated with prophylactic cranial irradiation (PCI) or not:		From 16-Sep-2	25	No	n/a	Yes	Agreed	No	30-Dec-25
		following platinum-based chemoradiotherapy where the following criteria have been met:	- yes, the patient has received PCI or - no, the patient has not been treated with PCI									
		citeria nave been met.	6. Treatment with durvalumab maintenance monotherapy will continue until disease progression or symptomatic deterioration or unacceptable toxicity or withdrawal of patient consent or for a maximum of 2 calendar years, whichever occurs first.									
			7. The patient will start his/her first treatment with durvalumab within 42 days from the last day of the final cycle of chemotherapy (e.g. C4D21) or the last day of radiotherapy, whichever occurs later.									
			8. The patient has a current ECOG performance status of 0 or 1.									
			9. The patient has no symptomatically active brain metastases or leptomeningeal metastases.									
			10. The patient has had no prior treatment with anti-PD-L1/PD-1 therapy for small cell lung cancer, unless this was received for this indication via a company early access program and all treatment criteria on this form are fulfilled.									
			11. When a treatment break of more than 12 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment. This must be approved before durvalumab is re-commenced									
			12. Durvalumab will be otherwise used as set out in its Summary of Product Characteristics (SPC).									

				Available	to new p	atients				Interim Funding	CDF	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Transition Funding agreed by manufacturer (Agreed, Rejected, Pending)	Eligible for Interim Funding (Yes, No, Not currently applicable (NCA))	agreed by manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Managed Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
ELR1	Eiranatamab	For the treatment of relapsed or refractory myeloma in adult patients who have relapsed or are refractory to their last anti-myeloma regimen AND have received at least 3 prior lines of systemic therapies which must have included at least one proteasome inhibitor, at least one immune-modulatory agent and at least one anti-CD38 antibody where the following criteria have been met:	1. This papilication for estimationable monothicapy is tooth being made by and the first cycle of systemic anti-cancer therapy with eiranatamab will be prescribed by a consultant specialist population for instead accredited in the use of systemic anti-cancer therapy.  2. The patient is an adult with a proven diagnosis of multiple myeloma.  Note: patients with amyloidosis or PDEMS syndrome are not eligible for efranatamab.  3. The prescriming inclinaria understands that the relavatumb is not flueded for amyloidosis patients (with the exception of patients who have a proven diagnosis of myeloma with an associated diagnosis of amyloidosis) and that NHS funding for efranatamab is only for the relapsed or refractory myeloma indication in the specific indication recommended by NHCE.  Please tick the relevant box below:  1. This patient has been proving the proving a proven diagnosis of primary amyloidosis or this patient has a proven diagnosis of primary amyloidosis or this patient has a proven diagnosis of progressive myeloma with an associated diagnosis of amyloidosis and efranatamab is being prescribed for the myeloma (and all other treatment creteria on this form apply)  4. This patient has been previously treated with at least one proteasome inhibitor.  Please confirm how many different proteasome inhibitors have been used to treat this patient's myeloma:  1. Immunomodulatory agent or  2. or more different proteasome inhibitors  2. or more different move many different immunomodulatory agents.  4. This patient has been previously treated with at least one immunomodulatory agents.  5. This patient has previously precived a pomalidomide-containing regimen or very composition of the patient by a pomalidomide containing regimen or very composition of the patient by a pomalidomide containing regimen or very composition from more more application of the patient by a pomalidomide containing regimen or very composition because of the patient by a pomalidomide containing regimen or very composition by a provision of the		m 21-Jun-24		No	n/a	Yes	Agreed	Yes	nca

				Availa	ble to new	patients		Transition	Eligible for	Interim Funding	CDF Managed	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Interim Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
ELR1	Elranatamab	For the treatment of relapsed or refractory myeloma in adult patients who have relapsed or are refractory to their last anti-myeloma regimen AND have received at least 3 prior lines of systemic therapies which must have included at least one proteasome inhibitor, at least one immune-modulatory agent and at least one aimt-CDB3 antibody where the following criteria have been met:	Please record below the ECOG performance status		From 21-Jun-	24	No	n/a	Yes	Agreed	Yes	nca
ENF1	Enfortumab vedotin in combination with pembrolizumab	Enfortumab vedotin with pembrolizumab for untreated, unresectable or metastatic urotheilal cancer, when platinum-based chemotherapy is suitable where the following criteria have been met:	1. This application has been made by and the first cycle of systemic anti-cancer therapy.  2. The patient has a histologically- or cytologically confirmed diagnosis of unresectable or metastatic urothelial cancer (i.e., cancer of the bladder, renal pelvis, ureter, or urethra). Patients with squamous or sarcomatoid differentiation or mixed cell types are eligible.  3. In respect of his/her treatment for unresectable/advanced disease and at the time of starting enfortumab vedotin & pembrolizumab, the patient is/was treatment-naïve to systemic therapy.  4. In the absence of enfortumab vedotin & pembrolizumab the patient would have been deemed eligible for treatment with displatin or carboplatin-based chemotherapy.  5. The patient does not have ongoing sensory or motor neuropathy of grade 2 or higher  6. At the time of commencing pembrolizumab the patient has/had not received prior treatment with any of the following in respect of their urothelial cancer: anti-PD-1, an		From 21-Aug-	25	No	n/a	Yes	Agreed	No	10-Dec-25

				Availa	able to new	patients		Transition	Eligible for	Interim Funding agreed by	CDF Managed	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)		manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
ENTIa_v1.1	Entrectinib	ENT1b which requires information as to this RECIST response assessment must then be completed for continuation of funding for entrectinib beyond the initial 12 week period otherwise the dispensing	1. This application is made by and the first cycle of systemic anti-cancer therapy with entrectinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient is aged 12 years or older. Entrectinib is only licensed in those aged 12 and above. If the patient is aged under 12 years, larotrectinib is licensed in this age group and can be accessed vide from AETA.  3. This patient has a proven histological diagnosis of a malignant solid tumour (ie a carcinoma or a sarcoma or melanoma or a brain or spinal cord tumour) and does NOT have a leakasam or a hymphomaor or myelona.  Please state below the site of origin of the patient's cancer and its specific histological type.  4. This patient has disease that is locally advanced or metastatic or would require surgical resection likely to result in severe morbidity. Please enter below the type of disease that is being retarded:  1. locally advanced disease for which systemic therapy has been indicated or "metastatic diseases or or which surgical resection is likely to result in severe morbidity.  5. This patient has no satisfactory systemic therapy options. A satisfactory systemic treatment option is defined as one which is funded by NHS England for the disease and indication in question, by claim the adjacent yet by col. rooffirm that the patient has already been treated with all the systemic therapy options funded by NHS England for the disease in question. As part of the evidence that NHC and NHS England with to see at the NHC re-appraisal of entrectinib in NHTX gene fusion positive patients, data will be specifically analysed as to systemic therapy to cally advanced/metastatic disease, and patient has no satisfactory systemic therapy options, as described above.  1. In Special that is been used a first line therapy of locally advanced/metastatic disease, as the patient has no satisfactory systemic therapy options, as described above.  1. The patient has not previously received treatment		From 25-Jun-	20	No	n/a	Yes	Agreed	Yes	nca

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				Availa	able to new	patients		Transition	Eligible for	Interim Funding agreed by	CDF Managed	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Interim Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
ENTIb_v1.0	Entrectinib	This form ENT1b requires information as to the RECIST response assessment made at 10 weeks after initiation of entrectinib. In addition, form ENT1b must be completed for continuation of funding for entrectinib to occur beyond the initial 12 week period otherwise the dispensing Trust will not receive reimbursement for further entrectinib.  Note: the ENT1a form is for the initiation of treatment with entrectinib and is only for funding of the first TMELVE weeks of entrectinib treatment. A PET/CT/MR scan of index assessable/measureable disease and the brain must be done prior to	3. A RECIST radiological assessment has been made of any metastatic intra-cerebral or CNS disease at 10 weeks after the start of entrectinib and I have indicated the outcome of this RECIST assessment below. If the patient does not have any metastatic intra-cerebral disease, please indicate in the relevant box. If the patient has a primary cerebral tumour, the response assessment should be done in the above box.  - the patient does not have any metastatic intra-cerebral disease or - the patient has a primary brain tumour and the response assessment has been done in the above section of this form or - complete response in the brain/CNS or - partial response in the brain/CNS or - progressive disease in the brain/CNS or - progressive disease in the brain/CNS  Please indicate how many weeks there were between date of start of entrectinib and date of above CT/MR response assessment scan:  4. The current clinical decision to continue or discontinue treatment with entrectinib is as set out below: - the patient will continue treatment with entrectinib ie has so far achieved a complete response or a partial response or has stable disease or		From 25-Jun-:	ю	No	n/a	Yes	Agreed	Yes	nca

				Availab	ole to new pa	atients						
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Transition Funding agreed by manufacturer (Agreed, Rejected, Pending)	Eligible for Interim Funding (Yes, No, Not currently applicable (NCA))	Interim Funding agreed by manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	CDF Managed Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
GLO2	<b>Giofitamab</b> In combination with gemcitabine and oxaliplatin	Glofitamab with gemcitabine and oxilipatin for treating relapsed or refractory diffuse large Br- cell lymphoma where the following criteria have been met:	Refractory/resistant DIBCL i.e. no response to first cycle of first line treatment.  - DIBCL that initially went into remission but subsequently relapsed.  6. The patient is not eligible for an autologous stem cell transplant.  7. The patient has not previously received a bispecific antibody treatment.  8. The patient has an ECOG performance status score of 0, 1 or 2.  9. Treatment with glofitamab, gemcitabine and oxaliplatin will be stopped at whichever of the following events occurs first:  - disease progression - unacceptable toxicity - withdrawal of patient consent - a total of eight cycles of glofitamab, gemcitabine and oxaliplatin plus four additional cycles of glofitamab monotherapy Note: once glofitamab is stopped after 12 cycles of treatment, it cannot be re-started.  10. When a treatment break of more than 6 weeks beyond the expected 3-weekly cycle length is needed, a treatment break approval form will be completed to restart treatment.	Fre	om 13-Nov-2	25	No	n/a	Yes	Agreed	No	nca
			11. Glofitamab with gemcitabine and oxaliplatin will be used as per the Summary of Product Characteristics (SPC).									

				Avai	lable to new	oatients						
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Transition Funding agreed by manufacturer (Agreed, Rejected, Pending)	Eligible for Interim Funding (Yes, No, Not currently applicable (NCA))	Interim Funding agreed by manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	CDF Managed Access Scheme (Yes, No, Not currently applicable (NCA))	
ISA1_v1.1	Isatuximab in combination with pomalidomide and dexamethasone	Isatuximab in combination with pomalidomide and dexamethasone for the	1. This application is being made by and the first cycle of systemic anti-cancer therapy with instruction continuation with pomalidomide and dexamethasone will be prescribed by a consultant specialists specifically trained and excredited in the use of systemic anti-cancer therapy.  2. The pattern has a diagnosis of multiple impeloma.  3. The pattern has reviewed 3 and only 3 prior lines of treatment and that the numbering of a line of treatment is in accordance with the international Myeloma Workshop Consensus recommendations for the uniform reporting of clinical trais (http://doi.org/10.1182/blood-2010-10.299487). A line of therapy is defined as one or more cycles of a planned restrient program. This may consist of one or more planned cycles of ingle-geart therapy or combination threngs, see well as a sequence of treatment as formed in a planned manner (e.g. induction chemotherapy/chemotherapies if followed by stem cell transplantation then maintenance is considered to be 1 line of therapy. A new line of therapy status when a planned general of observation off therapy is interrupted by a need for additional restrients of the seed of the program of the progr		From 15-Oct		No	n/a	Yes	Agreed	Yes	nca

				Availa	able to new p	atients						
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Transition Funding agreed by manufacturer (Agreed, Rejected, Pending)	Funding (Yes, No,	Interim Funding agreed by manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	CDF Managed Access Scheme (Yes, No, Not currently applicable (NCA))	
ISA2	is atuximab in combination with bortezomib, lenalidomide, and dexamethasone	treatment of UNTREATED multiple myeloma when a stem cell transplant is	1. This application is both being made by and the first cycle of systemic anti-cancer therapy with isatuximab in combination with bortezomib, lenalidomide and dexamethasone, will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has newly diagnosed multiple myeloma.  Note: this isatuximab indication is not funded for patients with primary amyloidosis.  Please confirm this by ticking the box below:  - this patient does not have a diagnosis of primary amyloidosis  3. The patient does not have a diagnosis of primary amyloidosis  3. The patient dos previously not received any systemic anti-cancer therapy for myeloma except for either an emergency use of a short course of corticosteroids before this treatment or the patient commenced induction therapy with the combination of daratumumab plus bortezomib, thalidomide and dexamethasone with the intention of proceeding to a stem cell transplant but despite responding to such treatment the patient is now ineligible for transplantation.  Please tick below which scenario applies to this patient:  - the patient has not received any prior systemic anti-cancer therapy  - the patient has only had an emergency use of a short course of corticosteroids  - the patient has only had an emergency use of a short course of corticosteroids  - the patient who have not responded to induction therapy with the combination of daratumumab plus bortezomib, thalidomide and dexamethasone with the intention of proceeding to a stem cell transplant to despite responding to such treatment the patient is now ineligible for transplantation.  Note: patients who have not responded to induction therapy with daratumumab plus bortezomib, thalidomide and dexamethasone are NOT allowed to switch to the isatuximab combination regimen outlined in this Blueteq form  4. The patient is ineligible for an autologous stem cell transplant.  5. Isatuximab will only be given in combination with bortezomib, lenalidomide and dexamethaso		rom 04-Sep	15	No	n/a	Yes	Agreed	No	23-Dec-25

				Availa	able to nev	patients		Transition	Eligible for	Interim Funding	CDF	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (bu notice o remova served	l No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Interim Funding (Yes, No, Not currently applicable (NCA))	agreed by manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Scheme (Yes, No, Not Currently	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
LARIa_v1.1	Larotrectinib	RECIST response on the repeated assessment	1. This application is made by and the first cycle of systemic anti-cancer therapy. 2. This pattern has a proven histological diagnosis of a malignant solid tumour (ie a carcinoma or a sarcma or melanoma or a brain or spiral cord tumour) and dees NOT have a leukacian or a hypinoma or myeloma. Please state the site of origin of the patient's cancer (NB if sarcoma, please enter sarcoma; if unknown primary, please state as such) and its specific histological type (eg for breast cancer dictal acromonia, localiz carcinoma, sceretory carcinoma etc. eg for lung cancer squamous NSCLC etc. eg for sarcoma: fibrosarcoma, osteoarcoma, gastrointestinal stromal tumour etc.)  3. This patient has desice that is locally advanced or metastatic or would require surgical resection likely to result in severe morbidity.  Please enter the type of disease that is being treated:  1. This patient has desice that is being treated:  1. This patient has disease that is locally advanced or metastatic or would require surgical resection likely to result in severe morbidity.  2. This patient has one with which surgical resection is likely to result in severe morbidity. Please state the type of surgical resection which would otherwise have been needed and restatic disease of work with surgical resection is likely to result in severe morbidity.  4. This patient has no satisfactory systemic therapy options. A satisfactory systemic treatment option is defined as one which is funded by NHS regiland for the disease in question.  4. This patient has no satisfactory systemic therapy options. A satisfactory systemic treatment option is defined as one which is funded by NHS England for the disease in question.  4. This patient has no satisfactory systemic therapy options funded by NHS England for the disease in question.  5. This patient has no satisfactory systemic therapy options funded by NHS England for the disease in question.  5. This patient has no satisfactory systemic therapy options funded by NHS England for the disease in question.  5. This		From 21-Api		No	nca	Yes	Agreed	Yes	nca

				Availa	ble to new	patients		Transition	Eligible for	Interim Funding agreed by	CDF Managed	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)		manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
LARIb_v1.0	Larotrectinib	Larotrectinib response assessment and treatment continuation form in the treatment of patients who have solid tumours that have a neurotrophic tyrosine receptor kinase (NTRI) gene fusion AND disease which is locally advanced or metastatic or for which surgical resection is likely to result in severe morbidity AND who have no satisfactory treatment options.  This form LAR1b requires information as to the REGST response assessment made at 10 weeks after initiation of larotrectinib. In addition, form LAR1b must be completed for continuation of funding for larotrectinib to occur beyond the initial 12 week period otherwise the dispensing Trust will not receive reimbursement for further larotrectinib.  Note: the LAR1a form is for the initiation of treatment with larotrectinib and is only for funding of larotrectinib and index assessable/measureable disease and index assessable/measureable/disease and the brain must be done prior to commencing larotrectinib and repeated at 10 weeks after the start of treatment (if not indicated before 10 weeks on account of assessing risk of disease progression).	Should be done in the above box.  - the patient does not have any metastatic intracerebral disease or - the patient does not have any metastatic intracerebral disease or - the patient has a primary brain tumour and the response assessment has been done in the above section of this form or - complete response in the brain/CNS or - stable disease in the brain/CNS or - stable disease in the brain/CNS or - stable disease in the brain/CNS - progressive disease in the brain/CNS  Please indicate how many weeks there were between date of start of larotrectinib and date of above CT/MR response assessment scan.  4. The current clinical decision to continue or discontinue treatment with larotrectinib is as set out below: - the patient will continue treatment with larotrectinib ie has so far achieved a complete response or a partial response or has stable disease or - the patient will discontinue or has discontinued treatment with larotrectinib on account of progressive disease or - the patient will discontinue or has discontinued treatment with larotrectinib on account of unacceptable toxicity  Note: RECIST-documented responses to larotrectinib in some patients can occur later than at 10 weeks and so a patient with stable disease would be expected to continue larotrectinib as long as the clinical assessment is that the patient signifies the early response rate.		From 21-Apr-	20	No	nca	Yes	Agreed	Yes	nca

				Availa	able to new	patients		Transition	Eligible for	Interim Funding	CDF Managed	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice o remova served)	NO	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed,	Interim Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
			1. This application for Iorlatinib is being made by and the first cycle of systemic anti-cancer therapy with Iorlatinib will be prescribed by a consultant specialist specifically trained and									
			accredited in the use of systemic anti-cancer therapy.  2. The patient has locally advanced or metastatic non-small cell lung cancer.									
			3. The patient has histological or cytological evidence of NSCLC that carries an anaplastic lymphoma kinase (ALK) rearrangement based on a validated test OR there is documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC AND there is an informative circulating free DNA test result confirming the presence of an activating anaplastic lymphoma kinase (ALK) rearrangement.  Please mark below on which basis the diagnosis of ALK positive NSCLC has been made in this patient:  - Histological or cytological evidence or  - Documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC and there is an informative circulating free DNA test result confirming the presence of an activating anaplastic lymphoma kinase (ALK) rearrangement									
LOR2	Lorlatinib monotherapy	Lorlatinib monotherapy for anaplastic lymphoma kinase-positive advanced non- small cell lung cancer previously untreated with an ALK inhibitor where the following criteria have been met:	4. The patient has not previously received any ALK inhibitor for the advanced NSCLC indication unless 1st line treatment with alectinib, brigatinib, certinib or crizotinib has had to be stopped within 6 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or the patient was treated with adjuvant alectinib and had disease progression more than 6 months after completing treatment with adjuvant alectinib.  Please mark below which of the five scenarios applies to this patient: - the patient has never previously received an ALK inhibitor or - the patient has previously received alectinib as 1st line ALK-targeted therapy and this has had to be stopped within 6 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or - the patient has previously received brigatinib as 1st line ALK-targeted therapy and this has had to be stopped within 6 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or		From 07-Oct	-25	No	n/a	Yes	Agreed	No	19-Jan-25
			5. The patient is treatment-naïve to 1st line cytotoxic chemotherapy-containing systemic treatment for this locally advanced or metastatic NSCLC indication.  Note: the only previous cytotoxic treatment allowed for patients to be treated with 1st line lorlatinib is adjuvant or neoadjuvant chemotherapy or chemotherapy given concurrently with radiotherapy.									
			6. The patient has an ECOG performance status of 0 or 1 or 2.									
			7. The patient either has no known brain metastases or if the patient has brain metastases, these must be asymptomatic (but can be treated or untreated.)									
			8. The patient will be treated until loss of clinical benefit or excessive toxicity or patient choice to discontinue treatment, whichever is the sooner.									
			<ol> <li>When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, the prescribing clinician will complete a treatment break approval form to restart treatment.</li> </ol>									
			Testant reactivents.  10. The prescribing clinician is aware that:  a) none of brigatinib or ceritinib or crizotinib are to be used following disease progression on lorlatinib as there is no current clear evidence to support treatment with any of these agents after disease progression on lorlatinib  and, therefore									
			b) after disease progression on lorlatinib, no subsequent ALK inhibitor therapy is commissioned by NHS England									
			11. Lorlatinib will otherwise be used as set out in its Summary of Product Characteristics (SPC).									

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				Availab	ble to new	patients		Transition	Eligible for	Interim Funding agreed by		
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)		Interim Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Managed Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
			1. This application for maintenance niraparib is being made by and the first cycle of systemic anti-cancer therapy with niraparib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient has a proven histological diagnosis of predominantly high grade serous or high grade endometrioid or high grade clear cell ovarian, fallopian tube or primary peritoneal carcinoma.  Please enter below as to which is the predominant histology in this patient:  - high grade serous adenocarcinoma or  - high grade endometrioid adenocarcinoma or  - high grade clear cell carcinoma									
		Niraparib monotherapy as maintenance treatment in patients with high grade epithelial stage III or IV ovarian, fallopian tube or primary peritoneal carcinoma who are in response following platinum-based FIRST line chemotherapy AND who HAVE a deleterious or suspected deleterious BRCA germline and/or somatic BRCA mutation (INCE TAG73) where the following criteria	4. This patient HAS a documented deleterious or suspected deleterious BRCA 1 or BRCA 2 mutation(s).  Please enter below as to which deleterious or suspected deleterious BRCA mutation(s) the patient has:  - BRCA 1 mutation or  - BRCA 2 mutation or									
NIR3_v1.2	Niraparib	have been met:  There is a separate form NIR4 for use of niraparib monotherapy as maintenance treatment in patients with high grade epithelial stage III or IV ovarian, fallopian tube or primary pertioneal carcinoma who are in response following platinum-based FIRST line chemotherapy and who DO NOT HAVE a deleterious or suspected deleterious BRCA germline and/or somatic BRCA mutation	5. The patient has recently diagnosed FiGO stage III or IV ovarian, fallopian tube or primary peritoneal carcinoma and has just completed 1st line platinum-based chemotherapy.  Note: maintenance niraparb in this 1st line maintenance indication is not funded for patients with recently diagnosed and treated stage I-III clisease.  6. One of the following scenarios applies to the surgical management of the patient in relation to the stage of the disease:  - the patient has stage III disease and had an upfront attempt at optimal cytoreductive surgery and had no visible residual disease at the end of surgery or  - the patient has stage III disease and had an interval attempt at optimal cytoreductive surgery and had visible disease at the end of surgery or  - the patient has stage III disease and had an interval attempt at optimal cytoreductive surgery and had visible disease at the end of surgery or  - the patient has stage III disease and had an interval attempt at optimal cytoreductive surgery and had visible disease at the end of surgery or  - the patient has stage IV disease and had an upfront attempt at optimal cytoreductive surgery and had no visible disease at the end of surgery or  - the patient has stage IV disease and had an upfront attempt at optimal cytoreductive surgery and had visible disease at the end of surgery or  - the patient has stage IV disease and had an interval attempt at optimal cytoreductive surgery and had visible disease at the end of surgery or  - the patient has stage IV disease and had an interval attempt at optimal cytoreductive surgery and had on visible disease at the end of surgery or  - the patient has stage IV disease and had an interval attempt at optimal cytoreductive surgery and had ovisible disease at the end of surgery or  - the patient has stage IV disease and had an interval attempt at optimal cytoreductive surgery and had ovisible disease at the end of surgery or  - the patient has stage IV disease and had an interval attempt at optimal cytoreductive surgery and had ovisi	F	From 15-Jan-	i 15-Jan-21	No	nca	Yes	Agreed	Yes	nca
			Please indicate below whether bevacizumab was used in combination with the 1st line chemotherapy: - bevacizumab 7.5mg/Kg given in combination with platinum-based chemotherapy or - bevacizumab 1.5mg/Kg given in combination with platinum-based chemotherapy or - no bevacizumab used in combination with chemotherapy  Criteria continue over the page									

				Availa	able to new	patients		Transition	Eligible for	Interim Funding agreed by	CDF Managed	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
NIR3_v1.1 (CONT)	Niraparib	are in response following platinum-based FIRST line chemotherapy AND who HAVE a deleterious or suspected deleterious BRCA germline and/or somatic BRCA mutation (TA673) where the following criteria have been met:  There is a separate form NIR4 for use of niraparib monotherapy as maintenance treatment in patients with high grade epithelial stage III or IV ovarian, fallopian tube or primary peritoneal carcinoma who are in response following platinum-based	15. Niraparib is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment, whichever is the sooner.  Note: NICE heard evidence during the niraparib appraisal that clinicians would wish to discuss with patients in continued complete remission when it would be an appropriate time to discontinue maintenance niraparib therapy and that this time was likely to be after approximately 3 years of maintenance treatment.		From 15-Jan-	21	No	nca	Yes	Agreed	Yes	nca

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				Availa	ble to new p	patients		Transition	Eligible for	Interim Funding agreed by	CDF Managed	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Interim Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
NIR4	Niraparib	are in response following platinum-based FIRST line chemotherapy AND who DO NOT HAVE a deleterious or suspected deleterious BRCA germline and/or somatic BRCA mutation [NICE TAG73]  There is a separate form NIR3 for use of niraparib monotherapy as maintenance treatment in patients with high grade epithelial stage lill or IV ovarian, fallopian	Please indicate below whether bevacizumab was used in combination with the 1st line chemotherapy:		From 15-Jan-2	21	No	nca	Yes	Agreed	Yes	nca

				Availal	ble to new p	atients		Transition	Eligible for	Interim Funding agreed by	CDF Managed	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)		Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Interim Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
			10. The patient will commence maintenance niraparib monotherapy within 12 weeks from the date of the first day of the last cycle of 1st line chemotherapy unless the patient was previously entered into the company's early access scheme for maintenance niraparib after 1st line chemotherapy and all the other treatment criteria set out in this form are fulfilled.  11. The patient has not previously received any PARP inhibitor unless the patient received 1st line maintenance rucaparib which has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression.  Please mark below which scenario applies to this patient:									
		Niraparib monotherapy as maintenance treatment in patients with high grade epithelial stage III or IV ovarian, fallopian tube or primary peritoneal carcinoma who are in response following platinum-based	The patient has never previously received a PARP inhibitor  - the patient has never previously received a PARP inhibitor  - the patient has a positive status for homologous recombination deficient disease and received 1st line maintenance rucaparib which has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression.  - the patient has a positive status for homologous recombination deficient disease and received 1st line maintenance rucaparib which has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression.  - The patient has a negative status for homologous recombination deficient disease and received 1st line maintenance rucaparib which has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression.									
NIR4 (CONT)	Niraparib	FIRST line chemotherapy AND who DO NOT HAVE a deleterious or suspected	12. Nuraparils win to eve as monotinerapy.  13. Maintenance niraparils is not being administered concurrently with maintenance bevacizumab.  14. The patient has an ECOG performance status of either 0 or 1.  Note: a patient with a performance status of 2 or more is not eligible for niraparils  Note: a patient with a performance status of 2 or more is not eligible for niraparils  15. Niraparils to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment, whichever is the sooner.  Note: NICE heard evidence during the niraparil appraisal that clinicians would wish to discuss with patients in continued complete remission when it would be an appropriate time to discontinue maintenance niraparils therapy and that this time was likely to be after approximately 3 years of maintenance treatment.	ı	From 15-Jan-2	1	No	nca	Yes	Agreed	Yes	nca
		treatment in patients with high grade epithelial stage III or IV ovarian, fallopian tube or primary peritoneal carcinoma who are in response following platinum-based FIRST line chemotherapy and who HAVE a deleterious or suspected deleterious BRCA germiline and/or somatic BRCA mutation										
		ga	12. The prescribing clinician understands that the marketing authorisation for niraparib recommends that full blood counts are performed weekly for the 1st month of treatment with niraparib.  13. The prescribing clinician understands that the marketing authorisation for niraparib recommends that the patient's blood pressure is monitored weekly for the first 2 months of treatment, monthly for the 1st year of therapy and then periodically thereafter during drug treatment with niraparib.  13. A first formal medical review as to whether maintenance treatment with niraparib should continue or not will be scheduled to occur at least by the start of the third 4-weekly cycle of treatment.									
			20. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment, including as appropriate if the patient had an extended break on account of Covid-19.  21. Niraparib is to be otherwise used as set out in its Summary of Product Characteristics									

				Availa	ble to new	patients		Transition	Eligible for	Interim Funding agreed by	CDF Managed	
Blueteq Form ref:	Drug	Indication	Indication  Criteria for use  Yes  1. This application is being made by and that leucapheresis for and treatment with obecabtagene autoleucel (obecel) will be initiated by a consultant haematologist specifically trained and accredited in the use of systemic anti-cancer therapy and working in an accredited CAR-T cell treatment centre and who is a member of the National CAR-T Clinical Panel for adult acute lymphoblastic leukaemia and a member of the treating Trust's adult acute lymphoblastic leukaemia and CAR-T cell multidisciplinary teams.	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Interim Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
OBE01a	Obecabtagene autoleucel	patients aged 26 years and older where the following criteria have been met:  This form is for the approval of leucapheresis and manufacture of CAR-T cells. There is a second part to this form which relates to the subsequent infusion of CAR-T cells and this will be available after submission of the first bard. The second part of the form (OBEO1b) can only be completed as a continuation of this first part of the form (OBEO1b) and OBEO1b must be completed on infusion of	and accredited in the use of systemic anti-cancer therapy and working in an accredited CAR-T cell treatment centre and who is a member of the National CAR-T Clinical Panel for adult acute lymphoblastic leukaemia and a member of the treating Trust's adult acute lymphoblastic leukaemia and CAR-T cell multidisciplinary teams.  2. The patient has CD19 positive relapsed or refractory B lineage acute lymphoblastic leukaemia (ALL).  Please tick apportate box as to which type of ALL the patient has:  - Philadelphia chromosome negative ALL or  - Philadelphia chromosome negative ALL or		From 25-Nov	-25	No	n/a	Yes	Agreed	No	tbc

				Availal	ble to new p	oatients		Transition	Eligible for	Interim Funding agreed by	CDF Managed	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)		Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Interim Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
OBE01b		Obecabtagene autoleucel (obecel) for treating relapsed/refractory Philadelphia negative and positive B cell acute lymphobastic leukaemia in patients aged 26 years and older where the following criteria have been met:  This second form is to document the date of infusion of CAR T cell therapy and for registration of this infusion with NHS England so that the treating Trust is reinflumented for the cost of obecabtagene autoleucel (obecel). There is a first form for the approval of leucapheresis and manufacture of CAR T cells. This second form must use the same unique Blueteq identifier number generated when this patient was registered for leucapheresis and CAR T cell manufacture using the first form.	3. The patient has an ECOG performance status of 0 or 1.  Please mark in the box below the current performance status: - PS 0 or - PS 1  The patient has sufficient end organ function to tolerate treatment with obscrabtagene autoleure! (obscell)	F	From 25-Nov-2	25	No	n/a	Yes	Agreed	No	tbc

				Availat	ble to new p	atients		Transition	Eligible for	Interim Funding agreed by	CDF Managed	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Interim Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
SEL4	Selpercatinib	Selpercatinib as monotherapy for the 1st line treatment of adult patients with previously untreated advanced non-small cell lung cancer (NSCLC) exhibiting a RET gene fusion where the following criteria have been met:	1. This application for seliperation is being made by and the first cycle of systemic anti-cancer therapy with seliperation will be prescribed by a consultant specialist specifically trained and according of the therapy.  2. The patient has locally advanced or metastatic non-small cell lung cancer.  3. The patient has labridogically or optiogically confirmed diagnosis of non-small cell lung cancer.  9. The patient has labridogically or optiogically confirmed diagnosis of non-small cell lung cancer.  9. Flease mark which type of NSCLC applies to this patient:  9. **Construction**  1. **This patient**, NSCLC has been shown to hardour a RET gene fusion as determined on a tumour tissue biopsy or a plasma specimen (liquid biopsy) or both.  9. **Plasma specimen (liquid biopsy) or both tumour tissue biopsy or a plasma specimen (liquid biopsy) or both tumour tissue biopsy or a plasma specimen (liquid biopsy) or both tumour tissue biopsy or a plasma specimen (liquid biopsy) or both tumour tissue biopsy or a plasma specimen (liquid biopsy) or both tumour tissue biopsy or a plasma specimen (liquid biopsy) or both tumour tissue biopsy or a plasma specimen (liquid biopsy) or both tumour tissue biopsy or a plasma specimen (liquid biopsy) or both tumour tissue biopsy or a plasma specimen (liquid biopsy) or both tumour tissue biopsy or a plasma specimen (liquid biopsy) or both tumour tissue biopsy or a plasma specimen (liquid biopsy) or both tumour tissue biopsy or a plasma specimen (liquid biopsy) or both tumour tissue biopsy or a plasma specimen (liquid biopsy) or both tumour tissue biopsy or a plasma specimen (liquid biopsy) or both tumour tissue biopsy or a plasma specimen (liquid biopsy) or both tumour tissue biopsy or a plasma specimen (liquid biopsy) or both tumour tissue biopsy or a plasma specimen (liquid biopsy) or both tumour tissue biopsy or a plasma specimen (liquid biopsy) or both tumour tissue and plasma specimen (liquid biopsy) or both tumour tissue		From 22-Jun-2	3	No	n/a	Yes	Agreed	Yes	nca

				Availa	able to nev	/ patients		Transition	Eligible for	Interim Funding	CDF	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (bu notice o remova served)	f No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed,	Interim Funding (Yes, No, Not currently applicable (NCA))	agreed by manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Managed Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
SOT1_v1.2	Sotorasib	Sotorasib as monotherapy for the treatment of adult patients with advanced non-small cell lung cancer (NSCLC) exhibiting a KRAS-G12C mutation and who have been previously treated with at least 1 prior systemic therapy for advanced NSCLC where the following criteria have been met:	1. This application for sotrorable being made by and the first cycle of systemic anti-cancer therapy. 2. The patient has locally advanced or metastatic nors mail cell lung cancer. 3. The patient has socially advanced or metastatic nors mail cell lung cancer. 3. The patient has socially advanced or metastatic nors mail cell lung cancer that has been shown to exhibit a RRAS G12C mutation using a validated assay and determined on a tumour issue biopsy or a plasma specimen (liquid biopsy) or both.  Please mark which per of specimen soppositive for the presence of the RRAS G12C mutation.  - Lumour tissue biopsy only or plasma specimen (liquid biopsy) only or a both tumour tissue and plasma specimen.  - After prescribing cliquid biopsy) only or a both tumour tissue and plasma specimen.  - After prescribing clinical has completed below the status of the patient's lung cancer with respect to other actionable mutations is involved to be present and that all commissioned targeted therapies have been fully explored for this mutation.  - After prescribing clinical has completed below the status of the patient's lung cancer with respect to other actionable mutations is involved to be present or - the NSCLC has an RRAS G12C in the NSCLC bits and RRAS G12C mutation is not to be present and that all commissioned targeted threapies have been explored for this read of the NSCLC base of the NSC	f	From 03-Ma	эг-22	No	n/a	Yes	Agreed	Yes	nca

				Availab	ole to new	patients				Interim Funding	CDF	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Transition Funding agreed by manufacturer (Agreed, Rejected, Pending)	Eligible for Interim Funding (Yes, No, Not currently applicable (NCA))	agreed by manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Managed Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
TALQ1	Talquetamab monotherapy	For treating relapsed or refractory multiple myeloma after 3 or more treatments where the following criteria have been met:	1. This application is being made by, wind drugs prescribed by a consultant or service resident doctor specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient is an adult with provine relipsed or refractory multiple myelona.  3. The patient has been or record or the patient of the patient of the patient is an adult with provine relipsed or refractory multiple myelona.  3. The patient has that 3 or more lines of treatment, according to the definition below, which must include:  - an immunomodiatory drug - a protessome inhibitor and - an artico38 antibody  - an artico38 antibody  The protection of the patient patie		om 17-Nov	-25	No	n/a	Yes	Agreed	No	tbc

				Availa	ble to new	patients				Interim Funding	CDF	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Transition Funding agreed by manufacturer (Agreed, Rejected, Pending)	Eligible for Interim Funding (Yes, No, Not currently applicable (NCA))	agreed by manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Managed Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
TRAD1_v1.1	Trastuzumab deruxtecan	For treating over-expressed HER2 positive unresectable locally advanced or uncastatic breast cancer in patients who have received 2 or more anti-HER2 therapies and who have received trastuzumab emtansine in the advanced/metastatic disease setting where the following criteria have been met:	1. This application for trasturumab denuteran for the treatment of unreactable locally advanced or metastatic breast cancer is being made by and the first cycle of trasturumab denuteran with processors before a consisting specialists specialisty specialists specialists pecialists specialists specialists specialists and consistent of the sure of systemic anti-cancer interapt.  2. The patient has intrologically documented breast cancer which is MEEQ 3+ by immunohistochemistry and/or has a MEEQ amplification ratio of 22.0 by in situ hybridisation.  4. If this patient received a HEEQ-targeted necadjournit regimen and its on tanture.  Pease text which points applies to this patient:  - the patient was not treated with a HEEQ-targeted necadjournit regimen.  - The patient was not treated with a HEEQ-targeted necadjournit regimen which contained trasturumab and trasturumab.  - The patient was received a HEEQ-targeted adjournit regimen which contained trasturumab as the sole HEEQ-targeted agent.  - If the patient received a HEEQ-targeted adjournit regimen which contained trasturumab and trasturumab.  - The patient was retard with a HEEQ-targeted adjournit regimen which contained trasturumab and trasturumab.  - The patient was retard with a HEEQ-targeted adjournit regimen which contained trasturumab and trasturumab.  - The patient was retard with a HEEQ-targeted adjournit regimen which contained trasturumab and trasturumab.  - The patient was retard with a HEEQ-targeted adjournit regimen which contained trasturumab and trasturumab.  - The patient was retarded with a HEEQ-targeted adjournit regimen which contained trasturumab and trasturumab.  - The patient was retreated with a HEEQ-targeted adjournit regimen which contained trasturumab and trasturumab.  - The patient was retreated with a HEEQ-targeted adjournit regimen for locally abounced/metastatic disease which included both perturumab and trasturumab.  - The patient was retreated with a HEEQ-targeted regimen for locally abounced/metastatic disease which included by t		From 20-Apr	21	No	n/a	Yes	Agreed	Yes	nca

				Availal	ble to new	patients				Interim Funding	CDF	
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	n No	Transition Drug (Old CDF) Indication (Yes or No)	Transition Funding agreed by manufacturer (Agreed, Rejected, Pending)	Eligible for Interim Funding (Yes, No, Not currently applicable (NCA))	agreed by manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	Managed Access Scheme (Yes, No, Not currently applicable (NCA))	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
TRAD2_v1.1	Trastuzumab deruxtecan	unresectable locally advanced or	1. This application for traitburnable drivesticen for the treatment of unresextable locally advanced or metastatic breast cancer is being made by and the first cycle of treatburnable devanced in the use of systems cancer the region in the control treatment in the use of systems cancer the region of the properties of the control of the		From 20-Dec		No	n/a	Yes	Agreed	Yes	nca

				Availab	le to new p	patients		Transition	Eligible for	Interim Funding agreed by		
Blueteq Form ref:	Drug	Indication	Criteria for use	Yes	Yes (but notice of removal served)	No	Transition Drug (Old CDF) Indication (Yes or No)	Funding agreed by manufacturer (Agreed, Rejected, Pending)	Interim Funding (Yes, No, Not currently applicable (NCA))	manufacturer (Agreed, Rejected, Pending, Not currently applicable (NCA))	y Managed trer Access , Scheme l, (Yes, No, tot Not y currently le applicable	Expected Entry into Baseline Commissioning (Date if known or Not currently applicable (NCA))
			1. This application for venetoclax plus obinutuzumab is being made by and the first cycle of this systemic anti-cancer therapy will be prescribed by a consultant specialist specifically									
			trained and accredited in the use of systemic anti-cancer therapy.									
			2. The patient has been diagnosed with chronic lymphatic leukaemia (CLL) or small lymphocytic lymphoma (SLL).									
			3. The patient has been tested for 17p deletion and the result is negative.  4. The patient has been tested for TPS3 mutation and the result is negative.									
											Managed Access Scheme (Yes, No, Not currently applicable	
			5. The patient has symptomatic disease which requires systemic therapy.  6. The patient has not received any previous systemic therapy for CLL/SLL.									
			The patient has not received any previous systemic therapy for CLC/SCL.      The patient has a performance status of 0 or 1 or 2.      The patient has a performance status of 0 or 1 or 2.									
			7. The patient has a performance status of OOF TO 2.									
			8. In the absence of this venetoclax plus obinutuzumab treatment option, the patient would otherwise have been treated with the combination of fludarabine, cyclophosphamide and rituximab (FCR) or the combination of bendamustine and rituximab (BR).  Please record below as to which combination you would have treated the patient with in the absence of this CDF access to venetoclax plus obinutuzumab:  - FCR or  - BR									
		For the treatment of patients with previously untreated chronic lymphatic	9. Venetoclax will be given in combination with obinutuzumab and that the venetoclax dose titration schedule will only be commenced after the patient has received the first 3 doses of obinutuzumab in cycle 1 (on days 1±2, 8 and 15) i.e. the venetoclax dose titration schedule is planned to commence on cycle 1 day 22 and be completed on cycle 2 day 28.									
VEN7_v1.1	Venetoclax in combination with obinutuzumab	leukaemia in whom chemotherapy with the combinations of either FCR or BR would otherwise have been SUITABLE where the following criteria have been met:	10. All of the following for the prevention and treatment of tumour lysis syndrome: - that the patient has been prospectively assessed for the risk of the development of tumour lysis syndrome (TLS) with venetodax - that appropriate TLS risk mitigation strategies have been put in place as outlined in the updated venetodax Summary of Product Characteristics - that there is a robust system in place for measuring appropriate blood chemistries both at the specified timings of blood chemistries according to TLS risk status and at the venetodax dose levels described in Section 4.2 Table 3 of the Summary of Product Characteristics. See https://www.medicines.org.uk/emc/medicine/32650 or https://products.mhra.gov.uk/substance/FSNBTCOLAX - that there is a robust system in place for ensuring the rapid review in real time of these blood chemistry results by a senior clinician with experience in the management of TLS - that there is a robust system in place for the withholding of the next days dose of each scheduled dose escalation until the blood chemistry results have been confirmed as being satisfactory by a senior clinician	Fn	om 10-Nov-	20	No	n/a	Yes	Agreed	Yes	nca
			11. The patient has been assessed specifically for potential drug interactions with venetoclax.									
			1.1. The patient has been assessed specificary for potential originate actions with venerous.  1.2. The maximum treatment duration of venetoclax in this indication is until day 28 of the 12th cycle of treatment i.e. the maximum duration of venetoclax treatment is for 45 weeks,									
			12. The maximum treatment our unition of venetociax in this indication is until day 28 of the 12th repeat of the maximum duration of venetociax treatment is not 45 weeks, closely find from cycle 1 day 25 followed by 11 cycles of 4-weekly cycles of 9 venetociax in cycles 2-12.									
			Losissing of 1 week non-type: 1 way 25 non-week of 11-type 30									
			13. Venetokas is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment or for the maximum treatment duration of 12 cycles (as									
			14. Venetoclask is to be communicated united usasses progression of unacceptable toxicity of patient choice to stop treatment of the maximum treatment during the statement of the statement of the maximum treatment during the statement of the									
			15. A formal medical review as to whether treatment with venetoclax in combination with obinutuzumab should continue or not will be scheduled to occur at least by the end of the									
			first 8 weeks of treatment.									
			16. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment,									
			including as appropriate if the patient had an extended break on account of Covid-19.									
			17. Venetoclax and obinutuzumab will be otherwise used as set out in their respective Summary of Product Characteristics (SPCs).									

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### B. NICE approved and baseline funded drugs/indications from 1st April 2016

If no Blueteq approval criteria are set this is because this was not considered necessary at the time of approval. However Blueteq registration will be required for all cancer drugs moving from the CDF to baseline as a result of positive final NICE guidance from 7th December 2016.

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application for abemaciclib in combination with an aromatase inhibitor is made by and the first cycle of systemic anti-cancer therapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The nation has histologically or cyclologically documented pastrogen recentor positive and her-2 pagetive breast cancer.				Startea
ABEM1_v1.2	Abemaciclib (in combination with an aromatase inhibitor)	The treatment of previously untreated, hormone receptor-positive, HER2-negative, locally advanced or metastatic breast cancer where the following criteria have been met:	2. The patient has histologically or cytologically documented oestrogen receptor positive and her-2 negative breast cancer  3. The patient has had no prior treatment with a CDK 4/6 inhibitor unless either palbociclib or ribociclib has had to be stopped within 6 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or a CDK 4/6 inhibitor has been previously received as adjuvant therapy and treatment was completed without disease progression at least 12 months prior to the first diagnosis of recurrent or metastatic disease.  Please mark below which one of these 4 scenarios applies to this patient:  - no prior treatment with a CDK 4/6 inhibitor or  - previous treatment with a CDK 4/6 inhibitor palbociclib but treatment has had to be stopped within 6 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of progressive disease or  - previous treatment with the 1st line CDK4/6 inhibitor vith endocrine therapy in the adjuvant setting for high risk early breast cancer either via NHS England commissioned care or via a clinical trial and treatment with the CDK 4/6 inhibitor with endocrine therapy in the adjuvant setting for high risk early breast cancer either via NHS England commissioned care or via a clinical trial and treatment with the CDK 4/6 inhibitor was completed without disease progression at least 12 months prior to the first diagnosis of recurrent or metastatic disease  4. The patient has metastatic breast cancer or locally advanced breast cancer which is not amenable to curative treatment  5. The patient has male or is female and if female is either post-menopausal or if pre- or peri-menopausal has undergone ovarian ablation or suppression with LHRH agonist treatment  6. The patient has had no previous hormone therapy for locally advanced or metastatic disease i.e. is hormone therapy malve for locally advanced/metastatic breast cancer.  Note: previous hormone therapy of procally advanced or eletrozole whet	No	TA563	27-Feb-19	28-May-19
			10. When a treatment treatment treat or more than a weeks beyond the expected	-			
			systemic anti-cancer therapy.  2. The patient has histologically or cytologically documented oestrogen receptor positive and HER-2 negative breast cancer  3. The patient has histologically or cytologically documented oestrogen receptor positive and HER-2 negative breast cancer  4. The patient is male or is female and if female is either post-menopausal or if pre- or peri-menopausal has undergone ovarian ablation or suppression with LHRH agonist treatment  5. The patient has an ECOG performance status of 0 or 1 or 2  6. The patient has received previous endocrine therapy according to one of the three populations as set out below as these are the only groups for which there was evidence submitted to NICE for the use of abemaciclib plus fulvestrant. Please record which population the patient falls into:  - has progressive disease whilst still receiving adjuvant or neoadjuvant endocrine therapy for early breast cancer with no subsequent endocrine therapy received following disease progression or  - has progressive disease whithin 12 or less months of completing adjuvant endocrine therapy for early breast cancer with no subsequent endocrine therapy received following disease progression or  - has progressive disease on 1st line endocrine therapy for advanced/metastatic breast cancer with no subsequent endocrine therapy received following disease progression or				
ABEM2	Abemaciclib (in combination with fulvestrant)	The treatment of hormone receptor- positive, HER2-negative, locally advanced or metastatic breast cancer where the following criteria have been met:	7. The patient has had no prior treatment with a CDK 4/6 inhibitor unless either palbociciib (in combination with fulvestrant) or ribociciib (in combination with fulvestrant) has had to be stopped within 6 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or a CDK 4/6 inhibitor has been previously received as adjuvant therapy and treatment was completed without disease progression at least 12 months prior to the first diagnosis of recurrent or metastatic disease.  Please mark below which one of the 4 scenarios applies to this patient: - no prior treatment with a CDK 4/6 inhibitor or - previous treatment with the CDK 4/6 inhibitor or - previous treatment with the CDK 4/6 inhibitor palbocicib in combination with fulvestrant but treatment has had to be stopped within 6 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of progressive disease or - revious treatment with the CDK 4/6 inhibitor ribocicib in combination with fulvestrant but treatment has had to be stopped within 6 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of progressive disease or - revious treatment with the CDK 4/6 inhibitor with endocrine therapy in the adjuvant setting for high risk early breast cancer either via NHS England commissioned care or via a clinical trial and treatment with the CDK 4/6 inhibitor was completed without disease progression at least 12 months prior to the first diagnosis of recurrent or metastatic disease	No	TA725	15-Sep-21	14-Dec-21
			8. The patient has had no prior treatment with fulvestrant 9. The patient has had no prior treatment with everolimus 10. Abemaciclib will only be given in combination with fulvestrant 11. Treatment will continue until there is progressive disease or excessive toxicity or until the patient chooses to discontinue treatment, whichever is the sooner 12. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, the prescribing clinician will complete a treatment break approval form to restart treatment. 13. Abemaciclib and fulvestrant will be otherwise used as set out in its Summary of Product Characteristics (SPC)				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application for abemacicilib in combination with endocrine therapy is being made by and the first cycle of abemaciclib plus endocrine therapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has early breast cancer.  3. The patient has histologically or cytologically documented hormone receptor-positive and HER-2 negative breast cancer.  4. The patient has high risk early breast cancer as defined by having either 4 or more positive axillary lymph nodes or 1-3 positive axillary lymph nodes and a primary tumour size of ≥5cm and/or histologically grade 3 disease. Please mark in the box below which category applies to this patient:  2-4 positive axillary lymph nodes or  1-3 positive axillary lymph nodes and a primary tumour size ≥5cm or  1-3 positive axillary lymph nodes and a primary tumour size ≥5cm and histological grade 3 disease or  1-3 positive axillary lymph nodes and a primary tumour size ≥5cm and histological grade 3 disease  5. The patient has completed definitive locoregional therapy (surgery with or without radiotherapy).  6. The patient has completed any adjuvant or neoadjuvant chemotherapy.				
АВЕМЗ	Abemacidib in combination with endocrine therapy	As adjuvant treatment for high-risk hormone receptor-positive and HER2- negative early breast cancer where the following criteria have been met:	Please mark in the box below the relevant treatment that the patient did or did not receive any adjuvant or recondityount chemotherapy or  - the patient did not receive any adjuvant or neoadjuvant chemotherapy or  - the patient received adjuvant chemotherapy only or  - the patient received neoadjuvant chemotherapy  7. The patient has received no more than 12 weeks of adjuvant endocrine therapy after completion of the last non-endocrine therapy (surgery or chemotherapy or radiotherapy).  8. The patient is male or female and if female, pre- or peri-menopausal and having adjuvant aromatase inhibitor therapy that the patient has undergone ovarian ablation or suppression with LHRH agonist treatment.  Please mark in the box below which category applies to this patient:  - female on adjuvant tamoxifien or  - post-menopausal female on adjuvant aromatase inhibitor therapy and LHRH agonist treatment/ovarian ablation or  - male	No	TA810	20-Jul-22	18-Oct-22
			9. The patient has an ECOG performance status of 0 or 1.  10. Abemaciclib is being given in combination with standard endocrine therapy.  11. The patient has had no prior treatment with a CDK 4/6 inhibitor unless the patient has suffered unacceptable toxicity on adjuvant ribociclib plus an aromatase inhibitor without any evidence of disease progression on treatment and fulfils the involved nodal and other criteria in criterion 4 above and the patient is transferring to treatment with adjuvant abemaciclib plus endocrine therapy. The treatment plan should be for a maximum CDK4/6 inhibitor treatment duration of 2 calendar years in all (time on ribociclib plus that on abemaciclib).  Please mark in the box below which scenario applies to this patient:  - the patient has suffered unacceptable toxicity on ribociclib plus an aromatase inhibitor without any evidence of disease progression and fulfils the involved nodal criteria in criterion 4 above and is transferring to treatment with adjuvant abemaciclib plus an endocrine therapy with any CDK4/6 inhibitor without any evidence of disease progression and fulfils the involved nodal criteria in criterion 4 above and is transferring to treatment with adjuvant abemaciclib plus an endocrine therapy with a treatment plan for a maximum CDK4/6 inhibitor treatment duration of 2 calendar years in all.  Note: patients who have commenced adjuvant ribociclib for disease stages which do not comply with criterion 4 are NOT eligible to switch to abemaciclib.  12. Treatment with abemaciclib will continue until there is progressive disease or excessive toxicity or until the patient chooses to discontinue treatment or for a maximum of 2 calendar years, whichever is the sooner. For patients switching from ribociclib, the maximum total CDK4/6 inhibitor treatment duration is for 2 calendar years (time on ribociclib plus time on abemaciclib).				
			13. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, the prescribing clinician will complete a treatment break approval form to restart treatment.  14. Abemaciclib will be otherwise used as set out in its Summary of Product Characteristics (SPC).				

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ABI1	Abiraterone	Abiraterone for treating metastatic hormone-relapsed prostate cancer before chemotherapy is indicated	1. This application is being made by and the first cycle of systemic anti-cancer therapy with abiraterone will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient either has a proven histological or cytological diagnosis of adenocarcinoma of the prostate or has presented with a clinical picture consistent with metastatic prostate cancer with both widespread bone metastases radiologically typical of prostate cancer and a serum PSA of ≥50 ng/mL.  3. This patient has hormone-relapsed (castrate-resistant) metastatic prostate cancer.  4. The patient has no or only mild symptoms after androgen deprivation therapy has failed.  5. Chemotherapy is not yet indicated.  6. One of the following applies to this patient as regards any previous use of 2nd generation receptor inhibitors (such as enzalutamide, darolutamide or apalutamide) or CYP17 enzyme inhibitors (such as abiraterone). Please enter below as to which scenario applies to this patient.  4. The patient has not been previously received any treatment with enzalutamide or darolutamide or apalutamide or abiraterone or  4. The patient has not been previously received any treatment with enzalutamide or darolutamide or apalutamide or abiraterone or  5. The patient has previously received any treatment with enzalutamide or darolutamide or apalutamide or abiraterone or  6. Abiraterone is to be given in combination with prednisolone  8. The patient has previously received and previously receive	Yes	TA387	27-Apr-16	26-Jul-16
ABI2	Abiraterone	For the treatment of patients with hormone-relapsed (castrate-resistant) metastatic prostate cancer with disease progression during or following treatment with docetaxel-containing chemotherapy where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with abiraterone will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient has a proven histological or cytological diagnosis of adenocarcinoma of the prostate or has presented with a clinical picture consistent with metastatic prostate cancer with both widespread bone metastases radiologically typical of prostate cancer and a serum PSA of 250 ng/ml.  3. This patient has hormone-relapsed (castration-resistant) metastatic prostate cancer.  4. The patient has been treated with docetaxel-containing chemotherapy and has progressed during or following treatment.  5. One of the following applies to this patient as regards any previous use of 2nd generation receptor inhibitors (such as enzalutamide, darolutamide or apalutamide) or CYP17 enzyme inhibitors (such as abiraterone). Please enter below as to which scenario applies to this patient:  - the patient has not previously received any treatment with enzalutamide or apalutamide or abiraterone or - the patient has previously received any treatment with enzalutamide or apalutamide or abiraterone or - the patient has previously received any treatment with enzalutamide or apalutamide or abiraterone or of disease progression  6. Abiraterone is to be given in combination with prednisolone  7. The patient has an ECOG performance status (PS) of 0 or 1 or 2.  8. Abiraterone is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment.  9. A formal medical review as to how abiraterone is being tolerated and whether treatment with enzalutamide should continue or not will be scheduled to occur at least by the start of the third 4-weekly cycle of treatment.  10. Where a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment, including indicating as appropriate if	Yes	TA259	27-Jun-12	25-Sep-12

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
Blueteq Form ref:	Abiraterone In combination with androgen deprivation therapy (ADT)	For the treatment of newly diagnosed high risk metastatic hormone-sensitive prostate cancer where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with abiraterone will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient either has a proven histological or cytological diagnosis of adenocarcinoma of the prostate or has presented with a clinical picture consistent with metastatic prostate cancer with both widespread bone metastases radiologically typical of prostate cancer and a serum PSA of at least 50 ng/mL.  3. The patient has newly diagnosed high risk metastatic prostate cancer that is hormone sensitive.  Note: patients who fulfil the clinical picture of metastatic prostate cancer as outlined in criterion 2 above but who do not have histological or cytological confirmation are considered to have high risk metastatic disease.  Note: an exception to this criterion is for the maintained supply of abiraterone following trial closure for patients who entered the STAMPEDE prostate cancer trial (ISRCTN78818544) and who continue to benefit from abiraterone treatment.  4. The patient has an ECOG performance status of either 0 or 1 or 2.  5. This patient has either not been treated with docetaxel and has currently received androgen deprivation therapy (ADT) for no longer than 3 months before starting an androgen receptor targeted agent or has been treated with docetaxel and has currently received and norm or than 9 months.  Please enter below as to which scenario applies to this patient  - the patient has not been treated with docetaxel and has currently received no more than 3 months of ADT before starting an androgen receptor targeted agent or  - the patient has not been treated with docetaxel and has currently received no more than 3 months of ADT before starting an androgen receptor targeted agent or  - the patient has not been treated with docetaxel and has currently received no more than 3 months of ADT before starting an androgen receptor targeted agent or  - the patient has not been t	drug/	with reference to NHSE Urgent Interim Commissioning Policy Proposition 2424	NICE	baseline funding
			7. The patient has not previously received any androgen receptor targeted agent unless the patient has received enzalutamide or apalutamide for newly diagnosed metastatic hormone-sensitive prostate cancer which had to be stopped because of dose-limiting toxicity in the clear absence of disease progression and the patient mas progressive disease following treatment with 2 years of ADT plus abiraterone with or without enzalutamide for high risk non-metastatic disease as part of the STAMPEDE trial (ISRCTN78818544) and did not progress whilst on such treatment and the patient meets all the other criteria listed on this form or the patient has represent disease following treatment and the patient meets all the other criteria listed on this form.  Please mark below which of these 4 clinical scenarios applies to this patient:  - the patient has not previously received any androgen receptor targeted agent  - the patient commenced enzalutamide/paralutamide/daralutamide which had to be stopped because of dose-limiting toxicity in the clear absence of disease progression and the patient meets all the other criteria listed here.  - the patient was treated with 2 years of ADT plus abiraterone with or without enzalutamide for high risk non-metastatic disease as part of the STAMPEDE trial and did not progress whilst on such treatment and the patient meets all the other criteria listed here  - the patient has high risk hormone sensitive prostate cancer treated with abiraterone as part of the STAMPEDE trial and has not progressed whilst on such treatment and the patient meets all the other criteria listed on this form  8. Abiraterone plus prednisolone is being given in combination with ADT.  9. Abiraterone is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment.  10. Where a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment, which MUST be approved before treatment is reco				

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ACA1_v1.2	Acalabrutinib monotherapy	For the treatment of patients with previously untreated chronic lymphatic leukaemia which has a 17 p deletion or TP53 mutation where the following criteria have been met:	1. This application for acabibutuhib is being made by and the first cycle of this systemic anti-cancer therapy. 2. The patient has been diagnosed with chronic lymphatic leukaemia (CLL) or small lymphocytic lymphoma (SLL). 3. The patient has been diagnosed with chronic lymphatic leukaemia (CLL) or small lymphocytic lymphoma (SLL). 4. The patient has been tested for 17p deletion and for TPS3 mutation or look. Please indicate the result of these tests below: - positive for 17p deletion and negative for TPS3 mutation or negative for 17p deletion and negative for 17p deletion and negative for 17p deletion and patient for the result of these tests below: - positive for 17p deletion and negative for 17p deletion and 1	No	TA689	21-Арг-21	20-Jul-21

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ACA2_v1.4	<b>Acalabrutinib</b> monotherapy	For the treatment of patients with previously treated chronic lymphatic leukaemia where the following criteria have been met:	1. This application for acalabrutinib is being made by and the first cycle of this systemic anti-cancer therapy.  2. The patient has been previously diagnosed with chronic lymphatic leukaemia (CLL) or small lymphocytic lymphoma (SLL).  3. The patient has been tested for 17p deletion and for TP53 mutation and the results are as shown below: negative for both 17p deletion and rP53 mutation or negative for 17p deletion and negative for TP53 mutation or negative for 17p deletion and positive for TP53 mutation  4. The patient has seen previously treated with systemic therapy for CLL/SLL  6. The patient has seen previously treated with systemic therapy for CLL/SLL  6. The patient is treatment naïve to a Bruton's kinase inhibitor or the patient has been previously treated with systemic discendance of discase progression or the patient has previously been treated with the 1st line combination of ibrutinib plus venetoclax and was still in response on completion of treatment but has since relapsed and this application will be the first use of a BTK inhibitor since the 1st line combination of ibrutinib plus venetoclax.  Please mark which of the 4 scenarios below applies to this patient:  - the patient has not received any previously bear previously commenced anabrutinib for relapsed/refractory CLL/SLL and azanubrutinib has had to be stopped solely because of dose-limiting toxicity and in the clear absence of disease progression or - the patient has not received any previously commenced ibrutinib for relapsed/refractory CLL/SLL and zanubrutinib has had to be stopped solely because of dose-limiting toxicity and in the clear absence of disease progression or - the patient has previously commenced ibrutinib for relapsed/refractory CLL/SLL and zanubrutinib has had to be stopped solely because of dose-limiting toxicity and in the clear absence of disease progression or - the patient has previously commenced ibrutinib for relapsed/refractory CLL/SLL and zanubrutinib has had to be stopped solely because of dose-limiting toxicity	No	TA689	21-Apr-21	20-Jul-21
			7. The patient has an ECOG performance status of 0 or 1 or 2.  8. Use of acalabrutinib in this indication will be as monotherapy.  Note: AstraZeneca did not submit evidence to NICE for consideration of acalabrutinib in combination with an anti-CD20 monoclonal antibody in this indication.  9. The prescribing clinician is aware that whereas the bioavailability of acalabrutinib CAPSULES is reduced by co-administration of an antacid or a proton pump inhibitor, acalabrutinib TABLETS can be safely co-administered with gastric acid reducing agents such as proton pump inhibitors, H2-receptor antagonists and antacids (see acalabrutinib's Summary of Product Characteristics).  Note: this distinction between acalabrutinib capsules and tablets is also important as stocks of acalabrutinib capsules will no longer be available from mid November 2023; existing stocks of acalabrutinib capsules should be used as soon as possible. Acalabrutinib tablets are currently available.  10. Acalabrutinib is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment, whichever is the sooner.  Note: Patients entered into the NIHR STATIC trial (NIHR ref: 52879) may be randomised to receive intermittent treatment as part of the trial protocol  11. A formal medical review as to whether treatment with acalabrutinib should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment.  12. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, a treatment break approval form will be completed to restart treatment.				
ACA3_v1.3	Acalabrutinib monotherapy	For the treatment of patients with previously untreated chronic lymphatic leukaemia which does not have a 17p deletion or a TP53 mutation and in whom chemotherapy with FCR or Bit is unsuitable where the following criteria have been met:	13. Acaibarturin will be otherwise used as set out in its Summary of Product Characteristics (SPC).  1. This application for acaibarutinib is being made by and the first cycle of this systemic anti-cancer therapy with acalabrutinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has been diagnosed with chronic lymphatic leukaemia (CLL) or small lymphocytic lymphoma (SLL).  3. The patient has been tested for 17p deletion and the result is negative.  4. The patient has been tested for 17p deletion and the result is negative.  5. The patient has been tested for 17p deletion and the result is negative.  6. The patient has been tested for 17p deletion and the result is negative.  7. The patient has been tested for 17p deletion and the result is negative.  8. The patient has been tested for 17p deletion and the result is negative.  8. The patient has been tested for 17p deletion and the result is negative.  9. The patient has been tested for 17p deletion and the result is negative.  9. The patient has not received any submission to NICE for the assessment of clinical and cost effectiveness of 1st line acalabrutinib in patients suitable for chemotherapy and hence NICE was unable to make a recommendation for this patient population.  9. The patient has not received any previous systemic therapy for CLL/SLL unless 1st line acalabrutinib was previously commenced via an AstraZeneca early access scheme or the patient commenced 1st line acalabrutinib and the zanubrutinib has had to be stopped solely because of dose-limiting toxicity and in the clear absence of disease progression.  9. The patient previously commenced 1st line acalabrutinib and the ranubrutinib has had to be stopped solely because of dose-limiting toxicity and in the clear absence of disease progression.  9. The patient previously commenced 1st line acalabrutinib and the anabrutinib has had to be stopped solely because of dose-limiting toxicity and in the clear abse	No	TA689	21-Apr-21	20-Jul-21

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ALE1_v1.5	Alectinib	For anaplastic lymphoma kinase-positive advanced non-small cell lung cancer previously untreated with an ALK inhibito where the following criteria are met:	1. This application for alectrinib is being made by and the first cycle of systemic anti-cancer therapy, with alectrinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has locally advanced or metastatic non-small cell lung cancer.  3. The patient has biotological evidence of NSELC Pland Denies is an informative circulating free DNA test result confirming the presence of an activating anaplastic lymphoma kinase (ALX) rearrangement. Please mark below on which basis the diagnosis of ALX positive NSELC has been made in this patient:  Please mark below on which basis the diagnosis of ALX positive NSELC has been made in this patient:  Occumented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSELC and there is an informative circulating free DNA test result confirming the presence of an activating anaplastic lymphoma kinase (ALX) rearrangement.  4. patient has not previously received any ALX inhibitor for the advanced NSELC indication unless 1st line treatment with loriatinib, brigatinib, certifinib or crizotinib has had to be stopped within 6 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or the patient has never previously received an ALX inhibitor.  Please mark below which of the five scenarios applies to this patient:  1. the patient has never previously received an ALX inhibitor for the advanced herapy and this has had to be stopped within 6 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or the patient has previously received brigatinib as 1st line ALX-cargeted therapy and this has had to be stopped within 6 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or the patient has previously received brigatinib as 1st line ALX-cargeted therapy and this ha	indication	TA536	Guidance  08-Aug-18	_
			and c) after disease progression during treatment with adjuvant alectinib or within 6 months of completion of treatment with adjuvant alectinib, re-treatment with alectinib is not commissioned.  11. Alectinib will otherwise be used as set out in its Summary of Product Characteristics (SPC).				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ALE2	Alectinib	only lile non-small cell uning cancer whose tumours have an ALK gene rearrangement where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with adjuvant alectinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a histologically documented non-small cell lung cancer (NSCLC).  3. The patient has undergone a complete resection of the NSCLC with all surgical margins negative for tumour.  4. The pathological stage determined on this patient's surgical NSCLC specimen was a stage IIA or IIB or IIIA or N2 only IIIB tumour according to the UICC/AICC TNM 8th edition.  Please mark below which stage applies to this patient:  - stage IIA disease (T12 N 0)  - stage IIB disease (T13 N 10 or T12 N 1 or T12 N 1 or T2 N 1 or T3 N 10)  - stage IIB disease (T13 N 2 or T15 N 2 or T12 N 2 or T2 N 2 or T3 N 10 or T4 N 1)  - stage IIB disease (T13 N 2 or T15 N 2 or T12 N 2 or T2 N 2 or T3 N 10 or T4 N 1)  - Stage IIB disease (T13 N 2 or T15 N 2 or T15 N 2 or T4 N 2)  5. The patient's NSCLC has been documented on the tumour specimen (biopsy or surgical specimen) as exhibiting an anaplastic lymphoma kinase (ALK) gene arrangement.  6. The patient did not receive any pre-operative systemic therapy (cytotoxic chemotherapy, immunotherapy, ALK-targeted tyrosine kinase inhibitors) for the NSCLC.  7. The patient did not receive any pre-operative or post-operative radiation therapy for the NSCLC.  8. No more than 12 weeks have elapsed since surgery  9. The patient has an ECOG performance status (PS) of 0 or 1.  11. The patient has an ECOG performance status (PS) of 0 or 1.  12. Alectinib will be administered as monotherapy.  13. The patient will be treated with alectinib for whichever is the sooner of: disease progression or unacceptable toxicity or withdrawal of patient consent or for a total treatment duration of 2 calendar years.  14. A formal medical review as to how alectinib is being tolerated and whether treatment with alectinib should continue or not will be scheduled to occur at least by	No	TA1014	13-Nov-24	11-Feb-25

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eteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application for alpelisib in combination with fulvestrant is being made by and the first cycle of alpelisib plus fulvestrant will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anticancer therapy.				
			2. The patient has histologically or cytologically documented hormone receptor positive and HER-2 negative breast cancer.				
			3. The patient's breast cancer has a PIK3CA mutation identified in a tumour or plasma specimen using a validated test.				
			Note: patients with an AKT1 or PTEN genomic alteration but without a PIK3CA genomic alteration are not eligible for alpelisib plus fulvestrant.				
			4. The patient has metastatic or locally advanced breast cancer which is not amenable to curative treatment.				
			5. The patient is male or female and if female is either post-menopausal or if pre- or peri-menopausal has undergone ovarian ablation or suppression with LHRH agonist treatment.				
			6. The patient has progressive disease after previous endocrine-based therapy.				
			7. The patient has been previously treated with an aromatase inhibitor. Please record in which places in the treatment pathway the patient had aromatase inhibitor therapy: - solely for early breast cancer or - solely for locally advanced/metastatic breast cancer or - in both early and advanced breast cancer settings				
			8. The patient has been previously treated with a CDK4/6 inhibitor. Please record in which places in the treatment pathway the patient had CDK4/6 inhibitor therapy: - solely for early breast cancer or - solely for locally advanced/metastatic breast cancer or - in both early and advanced breast cancer settings Note: the company submitted a case to NICE for consideration of clinical and cost effectiveness only in patients previously treated with a CDK4/6 inhibitor. This population is narrower than that in the marketing authorisation.				
		For treatment of hormone receptor-	9. The patient has had no prior treatment with fulvestrant for any indication unless this patient is switching from treatment with capivasertib plus fulvestrant due to toxicity (see criterion 10 below).	1			
ALP1	Alpelisib	positive, HER2-negative, locally advanced or metastatic breast cancer in patients	Note: the marketing authorisation of alpelisib states that the efficacy of alpelisib in combination with fulvestrant is not considered to be established in patients previously treated with fulvestrant.				
ALPI	in combination with fulvestrant	previously treated with a CDK4/6 inhibitor and an aromatase inhibitor where the following criteria have been met:		No b	18816	10-Aug-22	08-Nov-2
			Please record which scenario applies to this patient: - the patient has not previously received any treatment with a PIK3CA-targeted drug or - the patient has received previous treatment with capivasertib plus fulvestrant but such treatment with capivasertib plus fulvestrant but such treatment with capivasertib plus fulvestrant but such treatment of its start solely as a consequence of excessive toxicity and in the clear absence of disease progression and all other treatment criteria on this form apply				
			11. The patient has an ECOG performance status of 0 or 1.				
			12. Alpelisib will only be given in combination with fulvestrant.				
			13. Treatment with alpelisib will continue until there is progressive disease or excessive toxicity or until the patient chooses to discontinue treatment, whichever is the sooner.				
			14. Because the absorption of alpelisib is affected by food, the patients will be advised to take alpelisib immediately after food and at approximately the same time each day.		TA816		
			15. The prescribing clinician is aware of the potentially serious side-effects of alpelisib (e.g. hyperglycaemia, cutaneous reactions, diarrhoea, and pneumonitis) and of the necessary alpelisib dose adjustments for these toxicities, as outlined in alpelisib's Summary of Product Characteristics.				
			16. The prescribing clinician is aware that patients with a diagnosis of diabetes mellitus require a treatment consultation with a diabetic specialist or a healthcare professional experienced in the management of hyperglycaemia prior to the start of treatment with alpelisib.				
			17. Should the patient develop hyperglycaemia, a consultation with a healthcare professional experienced in the management of hyperglycaemia should be considered for all non-diabetic patients and is recommended for those patients who are any of the following: pre-diabetic or in those with a fasting blood glucose level >250mg/dL or >13.9 mmol/L or those have a BMI ≥30 or those of age ≥75 years.				
			18. The prescribing clinician is aware of the potential drug interactions between alpelisib and human Breast Cancer Resistance protein (BCRP) inhibitors and various cytochrome P450 enzyme systems, as outlined in alpelisib's Summary of Product Characteristics.				
			19. When a treatment break of up to 6 weeks beyond the expected 4-weekly cycle length is needed, the prescribing clinician will complete a treatment break approval form to restart treatment, including as appropriate if the patient had an extended break on account of Covid-19.				
			20. Alpelisib and fulvestrant will be otherwise used as set out in their respective Summaries of Product Characteristics (SPCs).				

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with apalutamide will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient has a proven histological or cytological diagnosis of adenocarcinoma of the prostate without neuroendocrine differentiation or features of a small cell carcinoma.  3. This patient has non-metastatic prostate cancer as defined by recent imaging with conventional imaging with both a whole body isotope bone scan and a CT/MR scan of the chest, abdomen and pelvis.  Note: patients with the sole abnormality of pelvic lymph nodes measuring <2cm in short axis diameter and which are below the aortic bifurcation are eligible for apalutamide in this indication.				
APA1	Apalutamide in combination with androgen deprivation therapy (ADT)	For the treatment of non-metastatic hormone-resistant (castration-resistant) prostate cancer in patients who are at high risk of developing metastatic disease where the following criteria have been met:	4. The patient has hormone-resistant (castrate-resistant) disease as defined by 3 rising PSA levels (after the nadir PSA level) and taken at least 1 week apart during androgen deprivation therapy.  5. The patient's serum testosterone level is <1.7mmol/L on gonadotrophin releasing hormone agonist/antagonist therapy or after bilateral orchidectomy.  6. The current PSA level is 22mg/ml.  7. The patient is at high risk of developing metastatic disease as defined by a PSA doubling time of \$10 months during continuous ADT.  Please document the actual PSA doubling time in the box below:  8. The patient has an ECOG performance status of either 0 or 1 or 2.  9. The patient has an ECOG performance status of either 0 or 1 or 2.  9. The patient has not previously received any 2nd generation androgen receptor inhibitors (such as enzalutamide, darolutamide) or CYP17 enzyme inhibitors (such as abiraterone) unless the patient received darolutamide for non-metastatic hormone-resistant (castration-resistant) which had to be stopped because of dose-limiting toxicity in the clear absence of disease progression and the patient meets all the other criteria listed on this form.  Please mark below which of these 2 clinical scenarios applies to this patient:  - the patient has not previously received any androgen receptor targeted agent  - the patient received darolutamide for non-metastatic hormone-resistant (castration-resistant) which had to be stopped because of dose-limiting toxicity in the clear absence of disease progression and the patient meets all the other criteria listed on this form  10. Apalutamide is being given only in combination with androgen deprivation therapy.	No	No TA740	28-Oct-21	26-Jan-22
			11. Apalutamide is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment.  12. A formal medical review as to how apalutamide is being tolerated and whether treatment with apalutamide should continue or not will be scheduled to occur at least by the start of the third 4-weekly cycle of treatment.  13. Where a treatment break of more than 6 weeks beyond the expected 4-week cycle length is needed, I will complete a treatment break approval form to restart treatment, including indicating as appropriate if the patient had an extended break because of COVID 19.  14. Apalutamide is to be otherwise used as set out in its Summary of Product Characteristics				
APA2	Apalutamide in combination with androgen deprivation therapy (ADT)	For the treatment of patients with newly diagnosed metastatic hormone-sensitive prostate cancer who are ineligible for chemotherapy with docetaed where the	1. This apatient has a proven histological or cytological diagnosis of adenocarcinoma of the prostate or has presented with a clinical picture consistent with metastatic prostate cancer with both widespread bone metastases radiologically typical of prostate cancer and a serum PSA of ≥50 ng/mL.  3. This patient has newly diagnosed metastatic prostate cancer that is hormone sensitive and has currently received androgen deprivation therapy (ADT) for no longer than 3 months before starting an androgen receptor targeted agent.  Please enter below as to which scenario applies to this patient:  - the patient has not yet received any ADT for metastatic prostate cancer or  - the patient has not yet received any ADT for metastatic prostate cancer or  - the patient has not received any upfront docetaxel chemotherapy for metastatic hormone sensitive prostate cancer.  5. The patient has an ECOS performance status (PS) of or 1 or 2.  6. The perscribed inclinate has assessed this patient's status as regards receiving upfront docetaxel and have concluded that the patient is ineligible for docetaxel on the grounds of either having significant comorbidities (i.e. the patient should not be treated with docetaxel) or the patient is lift for upfront docetaxel but after fully informed consent has chosen not to receive upfront docetaxel.  Please mark below which of these 3 clinical scenarios applies to this patient:  - the patient has significant comorbidities which preclude treatment with docetaxel (i.e. the patient should not be treated with docetaxel) or the patient should not be treated with docetaxel.  - the patient has significant comorbidities which preclude treatment with docetaxel (i.e. the patient should not be treated with docetaxel) or the patient of the patient of the search of the patient of the pa	No	TA741	28-Oct-21	26-Jan-22
			clear absence of disease progression and the patient meets all the other criteria listed here  - the patient was treated with 2 years of ADT plus abiraterone with or without enzalutamide for high risk non-metastatic disease as part of the STAMPEDE trial and did not progress whilst on such treatment and the patients meets all the other criteria listed here.  9. The patient has not previously received any apalutamide or any other androgen receptor targeted agent unless the patient has received apalutamide via a company early access scheme and the patient meets all the other criteria listed here.  10. Where a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment, which MUST be approved before treatment is recommenced.  11. Apalutamide is to be otherwise used as set out in its Summary of Product Characteristics.				

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ARS1	Arsenic trioxide	Arsenic trioxide for treating newly diagnosed low to intermediate risk acute promyelocytic leukaemia in ADULTS where all the following criteria are met:	1. An application is made by and the start of systemic anti-cancer therapy with arsenic trioxide will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The patient is an ADULT and has a confirmed diagnosis of acute promyelocytic leukaemia characterised by the presence of the t(15;17) translocation and/or the presence of the Pro-Myelocytic Leukaemia/Retinoic-Acid-Receptor-alpha (PML/RAR-alpha) gene  3. The patient is newly diagnosed with acute promyelocytic leukaemia  4. The patient has low to intermediate risk acute promyelocytic leukaemia (white cell count \$10 \times 10^9/L) and has not received any chemotherapy for this.  Patients with high risk acute promyelocytic leukaemia are not funded for treatment with arsenic trioxide  5. The patient will be treated with induction treatment of arsenic trioxide in combination with all-trans-retinoic acid (ATRA)  6. Induction treatment with arsenic trioxide will be continued until complete remission is achieved but if complete remission is not achieved by day 60, arsenic trioxide will be discontinued  7. As consolidation therapy, a maximum of 4 cycles of arsenic trioxide will be prescribed, each cycle being 4 weeks on treatment followed by 4 weeks off therapy  8. The dosing and schedule of administration of arsenic trioxide will be either in accordance with that described in the Summary of Product Characteristics (SPC) or that used in the UK NCRI AML17 trial as reported in Lancet Oncology 2015; 16:1295-1305.  If the AML17 dosing and schedule is used, hospital Trust policy regarding unlicensed treatments should be followed  9. The treating team is aware of the risk of and the treatment for  *APL differentiation syndrome  **O' Interval prolongation and the need for monitoring of electrolytes  **Liver toxicity  **Liver toxicity  **Do Arsenic trioxide is to be otherwise used as set out in its SPC	No	TA526	13-Jun-18	11-Sep-18
AR52	Arsenic trioxide	Arsenic trioxide for treating relapsed/refractory acute promyelocytic leukaemia in ADULTS where the following criteria are met:	1. An application is made by and the start of systemic anti-cancer therapy with arsenic trioxide will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. This patient is an ADULT and has a confirmed diagnosis of acute promyelocytic leukaemia characterised by the presence of the t(15:17) translocation and/or the presence of the Pro-Myelocytic Leukaemia/Retinoic-Acid-Receptor-alpha (PML/RAR-alpha) gene  3. The patient has acute promyelocytic leukaemia which is EITHER refractory to or relapsed after previous treatment which included a retinoid and chemotherapy OR has relapsed after a complete remission which lasted at least 2 years following previous arsenic trioxide and all-trans-retinoic acid treatment  4. The patient will be treated with induction and consolidation treatment of arsenic trioxide in combination with all-trans-retinoic acid (ATRA)  As combination therapy with ATRA is unlicensed in this relapsed/refractory setting, hospital Trust policy regarding unlicensed treatments should be followed  5. Induction treatment with arsenic trioxide will be continued until complete remission is achieved but if complete remission is not achieved by day 50 if the dosing and schedule is used as in the Summary of Product Characteristics or by day 60 if the UK NCRI AML 17 protocol is used (Lancet Oncology 2015; 16: 1295-1305), arsenic trioxide will be discontinued	No	TA526	13-Jun-18	11-Sep-18

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
AR53	Arsenic trioxide	Arsenic trioxide for treating newly diagnosed low to intermediate risk acute promyelocytic leukaemia in CHILDREN where the following criteria are met:	1. An application is made by and the start of systemic anti-cancer therapy with arsenic trioxide will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The patient is a CHILD and has a confirmed diagnosis of acute promyelocytic leukaemia characterised by the presence of the t[15;17] translocation and/or the presence of the Pro-Myelocytic Leukaemia/Retinoic-Acid-Receptor-alpha (PML/RAR-alpha) gene  3. The patient is newly diagnosed with acute promyelocytic leukaemia 4. The patient has low to intermediate risk acute promyelocytic leukaemia (white cell count ≤10 x 10°/L) and has not received any chemotherapy for this. Patients with high risk acute promyelocytic leukaemia are not funded for treatment with arsenic trioxide  5. The patient is mill be treated with induction treatment of arsenic trioxide in combination with all-trans-retinoic acid (ATRA)  6. Induction treatment with arsenic trioxide will be continued until complete remission is achieved but if complete remission is not achieved by day 60, arsenic trioxide will be discontinued  7. As consolidation therapy, a maximum of 4 cycles of arsenic trioxide will be prescribed, each cycle being 4 weeks on treatment followed by 4 weeks off therapy  8. The patient is a pre-pubescent or post-pubescent child and will be treated with edosing and schedule of administration of arsenic trioxide either in accordance with that described in the Summary of Product Characteristics (SPC) or that used in the UK NCRI AML17 trial as reported in Lancet Oncology 2015; 16: 1295-1305.  9. The use of arsenic trioxide has been discussed at a multi-disciplinary team (MDT) meeting which must include two consultants in the subspecialty with active and credible expertise in the relevant field of whom at least one must be a consultant paediatrician. The MDT should include a paediatric pharmacist and other professional groups appropriate to the disease area  10. The hospital Trust policy regarding unilicensed treatments	No	TA526	13-Jun-18	11-Sep-18
ARS4	Arsenic trioxide	Arsenic trioxide for treating relapsed/refractory acute promyelocytic leukaemia in CHILDREN where the following criteria have been met:	12. Arsenic trioxide is to be otherwise used as set out in its SPC  1. An application is made by and the start of systemic anti-cancer therapy with arsenic trioxide will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The patient is a CHILD and has a confirmed diagnosis of acute promyelocytic leukaemia characterised by the presence of the t(15;17) translocation and/or the presence of the Pro-Myelocytic Leukaemia/Retinoic-Acid-Receptor-alpha (PML/RAR-alpha) gene  3. The patient has acute promyelocytic leukaemia which is EITHER refractory to or relapsed after previous treatment which included a retinoid and chemotherapy OR has relapsed after a complete remission which lasted at least 2 years following previous arsenic trioxide and all-trans-retinoic acid treatment  4. The patient will be treated with induction and consolidation treatment of arsenic trioxide in combination with all-trans-retinoic acid (ATRA)  As combination therapy with ATRA is unlicensed in this relapsed/refractory setting, hospital Trust policy regarding unlicensed treatments should be followed  5. Induction treatment with arsenic trioxide will be continued until complete remission is achieved but if complete remission is not achieved by day 50 if the dosing and schedule is used as in the Summary of Product Characteristics or by day 80 if the UK NCRI AML 17 protocol is used (Lancet Oncology 2015; 16: 1295-1305), arsenic trioxide will be discontinued  6. As consolidation therapy, either the dosing and schedule in the Summary of Product Characteristics is used for a maximum of 3 vecks or the dosing and scheduling of the UK NCRI AML17 protocol (Lancet Oncology 2015; 16: 1295-1305), is used for a maximum of 4 cycles of arsenic trioxide, each cycle being 4 weeks on treatment followed by 4 weeks off therapy  7. The patient is a pre-pubescent or post-pubescent child and will be treated with the dosing and schedule of administration of arsenic trioxide either in accordance with	No	TA526	13-Jun-18	11-Sep-18

lueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. An application has been made by and the first cycle of arsenic trioxide will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
ARS5			2. The patient is aged >=18 years old and has a diagnosis of newly diagnosed high risk acute promyelocytic leukaemia (APML) as confirmed by:  • a white cell count >=10,000/µl (or 10 x 10 <sup>9</sup> /L) AND  • fusion of the PML/RARa gene (confirmed by fluorescence in situ hybridisation (FISH) analysis or PCR  3. The patient does not meet any of the following exclusion criteria:	-			
	Arsenic trioxide in combination with all-	Arsenic trioxide in combination with all- trans retinoic acid (ARTA) for the treatment of high-risk acute promyelocytic	<ul> <li>patient with isolated myeloid sarcoma but without evidence of APL by bone marrow or peripheral blood morphology</li> <li>patients with a pre-existing diagnosis of a prolonged QT syndrome, a history or presence of significant ventricular or atrial tachyarrhythmia, right bundle branch block plus left anterior hemiblock, bifascicular block</li> <li>patients on active diabysis for renal dysfunction</li> <li>female patients who are pregnant</li> </ul>	No	NHSE Policy:	N/A	05-Mar-25
	trans retinoic acid (ARTA)	leukaemia (>=18 years old) where the	hypersensitivity to arsenic trioxide or ATRA		URN2320	14/7	05 1110: 25
	,	following criteria are met:	4. The use of the arsenic trioxide will be discussed at a multi-disciplinary team (MDT) meeting which must include at least two haematology consultants.	I			
			5. The patient will receive the recommended dose and treatment regimen for arsenic trioxide as suggested in the NHS England Clinical Commissioning Policy.				
			6. The stopping / exit criteria have been explained and agreed with the patient and/or carer before the treatment is started and this has been documented in the patient records.				
			7. The Trust policy regarding unlicensed treatments has been followed.				
			NB. The use of arsenic trioxide in this indication is off-label, therefore Trust policy regarding unlicensed medicines should apply.  8. The patient has not previously received arsenic trioxide.  9. Arsenic trioxide will be otherwise used as set out in its Summary of Product Characteristics (SPC).	_			
			1. An application has been made by and the first cycle of arsenic trioxide will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			2. The patient is aged 12 months or older and has a diagnosis of newly diagnosed high risk acute promyelocytic leukaemia (APML) as confirmed by:				
			• a white cell count >=10.000/ul (or 10 x 10°/L) AND				
			• fusion of the PML/RARa gene (confirmed by fluorescence in situ hybridisation (FISH) analysis or PCR				
			3. The patient does not meet any of the following exclusion criteria:				
		Arsenic trioxide in combination with all-	patient with isolated myeloid sarcoma but without evidence of APL by bone marrow or peripheral blood morphology     patients with a pre-existing diagnosis of a prolonged QT syndrome, a history or presence of significant ventricular or atrial tachyarrhythmia, right bundle branch block plus left anterior hemiblock, bifascicular block     patients on active dialysis for renal dysfunction     female patients who are pregnant     hypersensitivity to arsenic trioxide or ATRA				
	Arsenic trioxide	trans retinoic acid (ARTA) for the	4. The use of the drug has been discussed at a specialised multidisciplinary team (MDT) meeting involving at least two paediatric haematological consultants who agree that continued treatment with arsenic trioxide is the most				
ARS6	in combination with all-	treatment of high-risk acute promyelocytic leukaemia (Children aged 12 months to	appropriate treatment plan. The MDT should also include a paediatric pharmacist and other professional groups appropriate to the disease area.	No	NHSE Policy: URN2320	N/A	05-Mar-25
	trans retinoic acid (ARTA)	<18 years old) where the following criteria have been met:	Patients should be discussed at a multidisciplinary team (MDT) prior to initiating treatment where time permits. However, in urgent cases where this is not possible, patients should be subsequently discussed at a local MDT meeting.		UNN2320		
			5. The patient will receive the recommended dose and treatment regimen for arsenic trioxide as suggested in the NHS England Clinical Commissioning Policy.				
			6. The stopping / exit criteria have been explained and agreed with the patient and/or carer before the treatment is started and this has been documented in the patient records.				
			7. The Trust policy regarding unlicensed treatments has been followed.				
			NB. The use of arsenic trioxide in this indication is off-label, therefore Trust policy regarding unlicensed medicines should apply.				
			8. The use of arsenic trioxide in this indication is being requested and administered in Principal Treatment Centres only.				
			9. The patient has not previously received arsenic trioxide.	1			
			10. Arsenic trioxide will be otherwise used as set out in its Summary of Product Characteristics (SPC).	1			
			11. Idarubicin chemotherapy will only be used during induction therapy and will follow the treatment regimen as suggested in the NHS England Clinical Commissioning Policy.	1			

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ASC1	Asciminib		1. This application for asciminib is being made by and the first cycle of systemic anti-cancer therapy with asciminib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has Philadelphia chromosome-positive chronic myeloid leukaemia (CML).  3. The CML remains in chronic phase.  4. The patient has received previous treatment with 2 or more TKIs for CML.  Please tick the appropriate option below as to the total number of different TKIs received by this patient:  2. 2 previous different TKIs  3. 2 previous different TKIS  5. The patient has been previously treated with ponatinib or not:  4. the patient has received treatment with ponatinib  5. The patient has received treatment with ponatinib  5. The patient has not received treatment with ponatinib  6. The last line of TKI therapy was either discontinued due to resistant disease in which case the T315I mutation test has been done and is negative or the last line of therapy was stopped due to patient intolerance of treatment in which case the previous T315I mutation test was negative.  4. The patient has not received previous different TKIs  5. The patient has not received previous T315I mutation test was negative.  5. The patient has not received previous T315I mutation test was negative.  6. The patient was intolerant of the last line of TKI therapy and the T315I mutation test was negative.  7. The patient was intolerant of the last line of TKI therapy and the previous T315I mutation test was negative.  8. The patient has not received prior treatment with asciminib unless the patient has started treatment via the Novartis compassionate use scheme and all other treatment criteria on this form are fulfilled.  8. The patient has not received prior treatment with asciminib via the EAMS scheme and all other treatment criteria on this form are fulfilled.  9. The patient started treatment with asciminib via the Novartis compassionate use scheme and all other treatment criteria on	No	TA813	03-Aug-22	02-Sep-22
			9. Asciminib will be given until the development of disease resistance or patient intolerance or withdrawal of patient consent.  10. The prescribing clinician understands that the daily dose of asciminib at the initiation of treatment for this indication is 80mg daily.				
			11. The prescribing clinician is aware of the potential drug interactions of asciminib with CYP3A4 inhibitors, CYP3A4 inducers, certain CYP3A4 substrates, CYP2C9 substrates and certain P-gp substrates.  12. The prescribing clinician is aware that asciminib absorption and bioavailability may be significantly reduced by concurrent administration with food (in particular high fat meals) and by some drugs (e.g. itraconazole) as described in asciminib's Summary of Product Characteristics).				
			13. A formal medical review as to how asciminib is being tolerated and whether treatment with asciminib should continue or not will be scheduled to occur at least by the end of the second 4-weekly cycle of treatment.				
			14. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment.				
			15. Asciminib will otherwise be used as set out in its Summary of Product Characteristics (SPC).	1			

lueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ATE1	Atezolizumab	The first line treatment of locally advanced or metastatic urothelial cancel in patients who are ineligible for cisplatin based chemotherapy and whose tumour have PD-L1 expression of 5% or more where all the following criteria are mett:	Please document the actual score for tumour inflitrating immune cell Pb-L1 expression:  1.1 The patient has not received prior treatment with an anti Pb-L1 anti-Pb-L1, anti-Pb-L2, anti-Pb-L2 anti-Pb	No	TA739	27-Oct-21	25-Jan-22
			12. The patient has no symptomatically active brain metastases or leptomeningeal metastases  13. Atecolizumab will be administered as monotherapy either subcutaneously at a dose of 1875mg every 3 weeks or intravenously at a dose of 1200mg every 3 weeks or 1680 mg every 4 weeks.  14. A formal medical review as to whether treatment with atezolizumab should continue or not will be scheduled to occur at least by the end of the third cycle of treatment.  15. The patient will be treated until loss of clinical benefit or excessive toxicity or patient choice to discontinue treatment, whichever is the sooner.				
			Note: there is no stopping rule for this indication.  16. When a treatment break of more than 3 months beyond the expected 3- or 4-weekly cycle is needed, a treatment break approval form will be completed to restart treatment.  17. Ateoroliumab will otherwise be used as set out in its Summary of Product Characteristics (SPC).	_			

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ATE2	Atezolizumab	Atezolizumab monotherapy for the treatment of PD-L1 positive or negative locally advanced or metastatic non-small cell lung cancer after chemotherapy where all the following criteria are met:	1. An application has been made by and the first cycle of systemic anti-cancer therapy.  2. The prescribing clinican's fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopaths, possibles and alkin notices.  3. The patient has a histologically-confirmed diagnosis of non-small cell lung cancer (quamuous or non-squamous).  4. The patient has personnel patient of the patient has a histologically-confirmed diagnosis of non-small cell lung cancer (quamuous or non-squamous).  5. PD-L1 stating with an approved and cellidated test to determine the Tumour Proportion Score (TS) has been attempted prior to this application and the result is set out below.  First, a please document the actual TFS below (if negative, record '07) or enter n\(^1\) of the the TFS cannot be documented and the reason why below.  First, a please and the patient is a proposed and cellidated test to determine the Tumour Proportion Score (TS) has been attempted prior to this application and the result is set out below.  First, a please and the patient is a proposed to the patient is a please document on the patient is a please of patient in the patient is a please	No	TA520	16-May-18	14-Aug-18

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. The application is made by and the first cycle of systemic anti-cancer therapy with atezolizumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy				
			2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for the immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, hepatitis and skin toxicities.				
			3. The patient has histologically or cytologically documented transitional cell carcinoma of the urothelial tract	1			
			4. The patient's disease is either locally advanced (ie T4b any N or any T N2-3 disease) or metastatic (any T any N M1 disease).				
			5. The patient has either not received previous adjuvant chemotherapy, neoadjuvant chemotherapy or chemo-radiotherapy, or if previously treated with platinum-based chemotherapy whether as adjuvant chemotherapy or as neoadjuvant chemotherapy or with chemo-radiotherapy, has relapsed =< 12 months since completing the platinum-based chemotherapy*.				
			* Patients meeting this criterion are eligible to be considered as previously treated for locally advanced/ metastatic disease (see below for criterion 6) but must satisfy all other criteria.				
			* Patients meeting this criterion are eligible to be considered as previously treated for locally advanced/ metastatic disease (and can answer "Yes" to criteria 6 below) but must satisfy all other criteria				
			6. There has been disease progression during or following previous platinum-based combination chemotherapy for inoperable locally advanced or metastatic urothelial cancer.	1			
			7. The patient has an ECOG performance status (PS) score of 0 or 1	1			
ATE3	Atezolizumab	Atezolizumab for locally advanced or metastatic urothelial cancer previously treated with platinum-based chemotherapy where all the following criteria are met:	8. The patient has not received prior treatment with an anti PD-1, anti-PD-12, anti-PD-12, anti-PD-13 or anti-cytotoxic T-lymphocyte-associated antigen-4 (CTL-4) antibody unless the patient completed or discontinued checkpoint inhibitor immunotherapy as part of adjuvant or neoadjuvant therapy without disease progression on treatment and at least 12 months elapsed between the date of last immunotherapy treatment and the date of first diagnosis of relapse with recurrent or metastatic disease.  Note: NHS England does not commission any re-treatment with checkpoint inhibitor therapy for patients who have discontinued or completed previous checkpoint inhibitor therapy for the locally advanced/metastatic indication.	No	TA525	13-Jun-18	13-Jul-18
			Please mark below if the patient has received previous checkpoint inhibitor therapy and in which setting:  - the patient has never received any immunotherapy for urothelial cancer. If so, please type 'n/a' in the 'Time gap' box below  - the patient has previously been treated with adjuvant immunotherapy for urothelial cancer and discontinued immunotherapy without disease progression and at least 12 months prior to the first diagnosis of disease relapse. Please document in the box below the time gap in months between completion of previous adjuvant immunotherapy and first diagnosis of disease relapse stable disease at the end of 1st line chemotherapy  - the patient has previously been treated with neoadjuvant treatment containing immunotherapy for urothelial cancer and discontinued immunotherapy without disease progression and at least 12 months prior to the first diagnosis of disease relapse. Please document in the box below the time gap in months between completion of previous neoadjuvant immunotherapy and first diagnosis of disease relapse  Time gap in months after completion of previous adjuvant or neoadjuvant recoding inhibitor immunotherapy and first diagnosis of disease relapse:				
			9. Atezolizumab will be administered as monotherapy either subcutaneously at a dose of 1875mg every 3 weeks or intravenously at a dose of 1200mg every 3 weeks or 1680 mg every 4 weeks.				
			10. A formal medical review as to whether treatment with atezolizumab should continue or not will be scheduled to occur at least by the end of the third cycle of treatment.				
			11. The patient is to be treated until disease progression and loss of clinical benefit or excessive toxicity or patient choice or <b>for a maximum treatment duration of 2 years of uninterrupted treatment</b> (ie a maximum of 35 administrations if given 3-weekly or a maximum of 26 administrations if given 4-weekly).				
			12. When treatment break of more than 3 months beyond the expected 3- or 4-weekly cycle length, a treatment break approval form will be completed.	1			
			13. The patient has no symptomatically active brain metastases or leptomeningeal metastases	-			
			14. Atezolizumab will otherwise be used as set out in its Summary of Product Characteristics (SPC)	1			

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
Blueteq Form ref:	Atezolizumab (in combination with bevacizumab, carboplatin and paclitaxel)	The first line treatment of adult patients with locally advanced or metastatic nonsquamous non-small cell lung cancer with a PD-L1 tumour proportion score of 0-49% and without EGFR and ALK mutations where the following criteria are met:	1. This application has been made by and the first cycle of systemic artificancer through with the combination of atezoidizumab, peractionab, carboplatin and pacificated will be prescribed by a consultant specialist specifically trained and accordibled in the use of systemic artificancer through.  2. As the peractioning clinican I am Infly waver of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-11 treatments including pneumonitis, collisis, nephritis, encountering the perial production of t	drug/ indication	TA TAS84	NICE	baseline funding
			13. The patient has no symptomatically active brain metastases or leptomeningeal metastases.  14. A formal medical review as to whether treatment with the combination of atezolizumab, bevacizumab, carboplatin and paclitaxel should continue or not will be scheduled to occur at least by the end of the first 6 weeks of treatment.  15. Where a treatment break of more than 12 weeks beyond the expected cycle length is needed, a treatment break form will be completed to restart treatment, including an indication as appropriate if the patient had an extended break because of COVID 19.  16. Atezolizumab and bevacizumab will be otherwise used as set out in their respective Summaries of Product Characteristics.				

Blueteq Form ref	: Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ATES	Atezolizumab (in combination with bevacizumab, carboplatin and paclitaxel)	The treatment of adult patients with EGFR or ALK or ROS1 or MET exon 14 or KRAS G12C or RET or BRAF mutation positive locally advanced or metastatic non-squamous non-small cell lung cancer after failure of appropriate targeted therapy where the following criteria are met:	1. The agriculture in twice made by a consultant specificit specifically trained and exceptional to a consultant specificit specifically trained and exceptional to a consultant specificit specifically trained and exceptional training or a consultant specific specifically trained and exceptional training or a consultant specificit specifically trained and exceptional training or a consultant specific specifically trained and exceptional training or a consultant specific specifically training or a consultant specific specifically confirmed diagnosis of non-squamous non-small cell lung cancer (MSCLC).  3. The patient has a shi biologically- or cyclopically- confirmed diagnosis of non-squamous non-small cell lung cancer (MSCLC).  4. The patient has a specific sp	No	TAS84	05-Jun-19	05-Jul-19

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ilueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with atezolizumab in combination with nab-paclitaxel will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies and hepatitis and skin toxicities.  3. The patient has a histologically- or cytologically-confirmed diagnosis of locally advanced and unresectable or metastatic breast cancer.				
ATE6_v1.1	Atezolizumab in combination with nab- pacilitaxei	For treating untreated PD-L1-positive, triple negative, unresectable, locally advanced or metastatic breast cancer for patients whose tumours express PD-L1 at a level of 1% or more where the following criteria have been met:	4. The patient's breast cancer has had receptor analysis performed and this is negative for all of the following: the HERZ receptor, oestrogen receptor and progesterone receptor i.e., the patient has triple negative disease.  5. The patient's tumour has been tested for PD-L1 expression and demonstrates PD-L1 expression of 1% or more by an approved and validated test.  Note: the measurement used for PD-L1 expression and demonstrates PD-L1 expression of low or more of the tumour area occupied by tumour cells, associated intra-tumoural and contiguous perl-tumoural desmoplastic stroma.  Please document the actual PD-L1 expression below:  PD-L1 expression below:  PD-L1 expression below:  PD-L1 expression below:  The patient has never had any prior treatment with anti-PD-L1/PD-L1 therapy for the breast cancer indication.  The patient has never had any prior treatment with anti-PD-L1/PD-L1 therapy for the breast cancer or the only previous anti-PD-L1/PD-L1 therapy.  Please mark below which of these clinical scenarios applies to this patient:  - the patient has never had any prior treatment with anti-PD-L1/PD-L1 therapy for the breast cancer or the only previous anti-PD-L1/PD-L1 therapy.  Please mark below which of these clinical scenarios applies to this patient:  - the patient has never had any prior treatment with anti-PD-L1/PD-L1 therapy for the breast cancer or  - the only previous anti-PD-L1/PD-L1 treatment that the patient has received was prior neoadjuvant and adjuvant therapy and there was no disease progression during such treatment and for at least 12 months after completion of anti-PD-L1/PD-L1 treatment that the patient has never had any prior treatment with anti-PD-L1/PD-L1 therapy.  Please document in the box below the time gap in months between completion of the previous neoadjuvant and adjuvant anti-PD-L1/PD-L1 immunotherapy and the first diagnosis of disease relapse. If the patient has never had such immunotherapy, please type n/s.  The patient is eligible for toxage non-vince page in months between co	No		01-Jul-20	31-Jul-20
АТЕ7	Atezolizumab in combination with carboplatin and etoposide	For the first-line treatment of adult patients with extensive-stage small cell lung cancer where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with atezolizumab in combination with carboplatin and etoposide will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, hepatitis and skin toxicity.  3. The patient has a histologically or cytologically determined diagnosis of small cell lung cancer.  5. The patient has been staged as having extensive stage small cell lung cancer.  5. The patient has not received previous systemic therapy for his/her extensive stage disease.  6. The patient has not received previous systemic therapy for his/her extensive stage disease.  6. The patient has an ECOG performance status score of 0 or 1.  7. The patient has an ECOG performance status score of 0 or 1.  7. The patient has an ECOG performance status score of 0 or 1.  8. On completion of 4 cycles of atezolizumab in combination with carboplatin and etoposide and in the absence of disease progression, reatment with atezolizumab maintenance monotherapy will continue until disease progression or symptomatic deterioration or unacceptable toxicity or withdrawal of patient consent, whichever occurs first.  9. Atezolizumab will be administered either subcutaneously at a dose of 1875mg every 3 weeks or intravenously at a dose of 1200mg every 3 weeks or 1680 mg every 4 weeks.  10. The patient has no symptomatically active brain metastases or leptomeningeal metastases.  11. The patient has had no prior treatment with anti-PD-L1/PD-1 therapy for small cell lung cancer.  12. A formal medical review as to how treatment with anti-PD-L1/PD-1 therapy for small cell lung cancer.  13. Where treatment break of more than 12 weeks beyond the expected 3-weekly or 4-weekly cycle length is needed,	No	TA638	01-Jul-20	31-Jul-20

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ATES	Atezolizumab in combination with bevacizumab	For the first-line systemic treatment of adult patients with locally advanced or metastatic and/or unresectable hepatocellular carcinoma where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy, with atezolizumab in combination with bevacizumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a diagnosis of hepatocellular carcinoma and that one of the following applies to the patient (please tick appropriate box below as to which option applies):  - either option 1 applies in which case the patient has a confirmed histological diagnosis of hepatocellular carcinoma (HCC) - or option 2 applies in which case a biopsy is deemed to be very high risk or technically not feasible in the patient and both the criteria a and b below are also both met: - as the decision not to biopsy has been made and documented by a specialist HCC multi-disciplinary team meeting - and be the tumour meets the non-invasive diagnostic criteria of HCC as set out below*.  It is expected that option 2 will only apply in exceptional circumstances.  Please mark below which of these 2 clinical senarios applies to this patient: - Option 1: the patient has a confirmed histological diagnosis of hepatocellular carcinoma or - Option 2: the patient has a confirmed histological diagnosis of hepatocellular carcinoma or - Option 2: the patient has a confirmed histological diagnosis of hepatocellular carcinoma or - Option 2: the patient has a confirmed histological diagnosis of hepatocellular carcinoma or - Option 2: the patient phase and the patient of the patient of the patient phase with washout in the portal venous or delayed phases). While one imaging technique is required for nodules beyond 1cm in diameter, a more conservative approach with 2 techniques is recommended in suboptimal settings.	No	TA666	16-Dec-20	15-Jan-21

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ATE9	Atezolizumab	of tumour cells or in at least 10% of	1. This application is being made by and the first cycle of systemic anti-cancer therapy with accelulament monotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a histologically- or cyclogically-confirmed diagnosis of non-mail cell lung cancer (squamous or non-squamous).  Passes made below with histology applies to this patient:	No	TA705	02-Jun-21	31-Aug-21

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ATE10	Atezolizumab	Atezolizumab monotherapy for adjuvant treatment after complete tumour resection in adult patients with UICC/AICC 8th edition stage IIB or III or 70 20 nml IIB non-small cell lung cancer and whose disease is all of the following: has Pol-1 expression on 250% of tumour cells, is not EGFR mutant or ALK-positive and has not progressed or recently completed adjuvant platinum-based chemotherapy where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with aljournal aterolisumab will be proscribed by a consultant specialist specifically trained and accordinate in the use of systemic anti-cancer therapy.  2. The preceding clinician is fully aware of the management of and the treatment modifications that may be required for immune-related alverse reactions due to anti-PD-L1 treatments including pneumonitis, collis, nephritis, educomospatibles, hepatitis and shift histories.  Please man below which historiesy applies to this patient:  - rans-spannous RSCLC - non-squamous RSCLC and the science of the patient of t	No	TA1071	19-Jun-25	21-Jul-25

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Blueteq Form ref	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ATE10	Atezolizumab	treatment after complete tumour resection in adult patients with UIC/AIC 5th edition stage IIB or IIIA or N2 only IIIB non-small cell lung cancer and whose disease is all of the following: has PD-L1 expression on 250% of tumour cells, is not EGFR mutant or ALK-positive and has not progressed on recently completed adjuvant platinum-based	16. Atezolizumab will be administered as monotherapy either subcutaneously at a dose of 1875mg every 3 weeks or intravenously at a dose of 1200mg every 3 weeks or 1680 mg every 4 weeks.  17. A formal medical review as to how atezolizumab is being tolerated and whether treatment with atezolizumab should continue or not will be scheduled to occur at least by the end of the second month of treatment.	No	TA1071	19-Jun-25	21-Jul-25

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
AVA1	Avapritinib monotherapy	For the treatment of aggressive systemic mastocytosis or aggressive systemic mastocytosis with an associated haematological neoplasm or mast cell leukaemia where the following criteria have been met:	1. This application for avapritinib monotherapy is being made by and the first cycle of systemic anti-cancer therapy with avapritinib monotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient is an adult and has a pathologically-confirmed diagnosis of aggressive systemic mastocytosis (ASM) or aggressive systemic mastocytosis with an associated haematological neoplasm (ASM-AHN) or mast cell leukaemia.  3. The patient has advanced disease and requires systemic therapy for this condition.  4. The patient has previously received systemic therapy for this condition or not.  Please mark below whether the patient has/has not previously received any previous systemic therapy for this condition  - yes, this patient has not received any previous systemic therapy for this condition  - yes, this patient has not received any previous systemic therapy for this condition  - yes, this patient has previously received midostaurin or not.  Please mark below whether the patient has previously received midostaurin or not.  1. The patient has not received previous midostaurin or not.  2. The patient has not received previous midostaurin  - yes, this patient has not received previous midostaurin  - yes, this patient has not received previous midostaurin  - yes, this patient has not received previous midostaurin  - yes, this patient has not received previous midostaurin  - yes, this patient has not received previous midostaurin  - yes, this patient has not received previous midostaurin  - yes, this patient has not received previous midostaurin  - yes, this patient has not received previous midostaurin  - yes, this patient has not received previous midostaurin  - yes, this patient has not received previous midostaurin  - yes, this patient has not received previous midostaurin  - yes, this patient has not received previous midostaurin  - yes, this patient has not received previous midostaurin  - yes, this patient has not received previous mid	No	TA1012	06-Nov-24	04-Feb-25

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. An application is made by and the first cycle of systemic anti-cancer therapy with avelumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy				
			2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for the immune-related adverse reactions due to anti-PD-11 treatments including pneumonitis, colitis, nephritis, endocrinopathies and hepatitis				
			3. The patient has a confirmed histological or cytological diagnosis of Merkel cell carcinoma				
			4. The patient has metastatic disease				
		The treatment of previously untreated	5. The patient is treatment naīve to any systemic anti-cancer therapy for Merkel cell carcinoma and in particular has not received any prior treatment with any anti-PD-1, anti-PD-11, anti-PD-12, anti-PD-12, anti-CD137 or anti-cytotoxic T-lymphocyte-associated antigen-4 [CTLA-4] antibody				
11/54		(with systemic therapy) metastatic Merkel	6. The patient has an ECOG performance status of either 0 or 1. Note: a patient with a performance status of 2 or more is not eligible for avelumab	1		24.4.24	20 1 1 24
AVE1	Avelumab	cell carcinoma where all the following	7. If the patient has brain metastases, then these have been treated and are stable	No	TA691	21-Apr-21	20-Jul-21
		criteria are met:	8. Avelumab is to be used as monotherapy only	- J			
			9. Avelumab is to be continued until loss of clinical benefit or unacceptable toxicity or patient choice to stop treatment. I also confirm that patients with radiological disease progression not associated with significant clinical deterioration (defined as no new or worsening symptoms and no change in performance status for greater than 2 weeks and no need for salvage therapy; all 3 conditions must apply) can continue treatment				
			10. A formal medical review as to whether treatment with avelumab should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment	1			
		11. Where a treatment break of more than 12 weeks beyond the expected cycle length of avelumab is needed, I will complete a treatment break approval form to restart treatment, including indicating as appropriate if the patient has an extended break because of COVID 19.	t				
			12. Avelumab will otherwise be used as set out in its Summary of Product Characteristics (SPC).	1			
			1. An application is made by and the first cycle of systemic anti-cancer therapy with avelumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy			-	
			2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for the immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies and hepatitis				
			3. The patient has a confirmed histological or cytological diagnosis of Merkel cell carcinoma	1			
			4. The patient has metastatic disease				
		The treatment of previously treated (with	5. I confirm that the patient has previously been treated with cytotoxic chemotherapy for metastatic Merkel cell carcinoma and has not received any prior treatment with any anti-PD-1, ant				
AVE2	Avelumab	systemic cytotoxic chemotherapy)	6. The patient has an ECOG performance status of either 0 or 1. Note: a patient with a performance status of 2 or more is not eligible for avelumab	No	TA517	11-Apr-18	10-Jul-18
		metastatic Merkel cell carcinoma where al the following criteria are met:	7. If the patient has brain metastases, then these have been treated and are stable				
		the following criteria are met:	8. Avelumab is to be used as monotherapy only	1			
			9. Avelumab is to be continued until loss of clinical benefit or unacceptable toxicity or patient choice to stop treatment. I also confirm that patients with radiological disease progression not associated with significant clinical deterioration (defined as no new or worsening symptoms and no change in performance status for greater than 2 weeks and no need for salvage therapy: all 3 conditions must apply) can continue treatment				
			10. A formal medical review as to whether treatment with avelumab should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment	1			
			11. Treatment breaks of up to 12 weeks beyond the expected cycle length of avelumab are allowed but solely to allow immune toxicities to settle				
			12. Avelumab will otherwise be used as set out in its Summary of Product Characteristics (SPC)	1		1	

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
AVE4_v1.0	Avelumab	Avelumab monotherapy for the maintenance treatment of adult patients with locally advanced or metastatic urothelial carcinoma who have just completed and not progressed on 1st lin platinum-containing combination chemotherapy where the following criteri have been met:	8. The patient will commence treatment with avelumab within 4 to 10 weeks of receiving the last dose of chemotherapy.	No	TA666	16-bec-20	15-Jan-21

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
AXI01a	Axicabtagene ciloleucel	treated with two or more lines of systemic therapy where the following criteria are met:  This form is for the approval of leucopheresis and manufacture of CAR-T cells. There is a second part to this form which relates to the subsequent infusion of CAR-T cells and this will be available after submission of the first part. The second part of the form (AXIOIa) can only be completed as a continuation of this first part of the form (AXIOIa) and must be completed on infusion of CAR-T cells otherwise the treating Trust will not be reimbursed for the cost of axicobtagene ciloleucel	-re-biopsy at second relapse has confirmed DLBL or PMBCL or -re-biopsy at second relapse was/is unsafe plus there is progressive disease at previously documented sites of active disease and the previous histology was DLBCL or PMBCL or -re-biopsy at second relapse has again confirmed transformed lymphoma (TFL, MZL, CLL, NLPHL) to DLBCL or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed FITLD of DLBCL type or -re-biopsy at second relapse has again confirmed F	Yes	TAS7Z	28-Feb-23	29-May-23

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
		Axicabtagene ciloleucel for treating relapsed/refractory diffuse large B-cell lymphoma (DLBCL), primary mediastinal B-cell lymphoma (DLBCL) and transformed lymphoma to DLBCL in patients previously treated with two or more lines of systemic therapy where the following criteria are met:  This form is for the approval of	12. The patient has an ECOG performance score of 0 or 1. Please enter below as to the patient's current ECOG performance status (PS): The ECOG performance status scale is as follows: PS 1 The patient is fully active and able to carry on all pre-disease performance without restriction PS 1 The patient is restricted in physically strenuous activity but is ambulatory and able to carry out work of a light or sedentary nature eg light house work, office work PS 2 The patient is restricted in physically strenuous activity but is ambulatory and able to carry out work of a light or sedentary nature eg light house work, office work PS 2 The patient is ambulatory and capable of all selfcare but unable to carry out any work activities and is up and about more than 50% of waking hours PS 3 The patient is capable of only limited selfcare and is confined to bed or chair more than 50% of waking hours PS 4 The patient is completely disabled, cannot carry out any selfcare and is totally confined to bed or chair The patient currently has a performance status of either - ECOG PS 0 or - ECOG PS 1				
AXI01a	Axicabtagene ciloleucel	leucapheresis and manufacture of CAR-T cells. There is a second part to this form which relates to the subsequent infusion of CAR-T cells and this will be available after submission of the first part. The second part of the form (AXIO1b) can only be completed as a continuation of this first part of the form (AXIO1a) and must be completed on infusion of CAR-T cells otherwise the treating Trust will not be reimbursed for the cost of axicobtagene ciloleucel	13. The patient has sufficient end organ function to tolerate treatment with CAR-T cell therapy.  14. The patient has either had no previous therapy with any genetically modified autologous or allogeneic T cell immunotherapy or the patient has been treated with doses of genetically modified autologous or allogeneic T cell immunotherapy within an abandoned dosing cohort in a first in human dose-escalation phase I clinical trial.  Please tick appropriate box as to which type of previous treatment the patient has had:  No previous therapy with any genetically modified autologous or allogeneic T cell immunotherapy or  - Previously treated with doses of genetically modified autologous or allogeneic T cell immunotherapy or  - Previously treated with doses of genetically modified autologous or allogeneic T cell immunotherapy within an abandoned dosing cohort in a first in human dose-escalation phase I clinical trial  15. Prior to infusion 2 doses of tocilizumab are available for use in this patient in the event of the development of cytokine release syndrome.  16. Axicabtagene ciloleucel-modified CAR-T cell therapy will otherwise be used as set out in its Summary of Product Characteristics (SPC).  17. Approval for the use of axicabtagene ciloleucel has been formally given by the National DLBCL/PMSCL/TFL CAR-T cell Clinical Panel.  Please state date of approval (DD/MM/YYYY)  18. Following national approval for use of axicabtagene ciloleucel there has been local CAR-T cell multidisciplinary team agreement that this patient continues to have the necessary fitness for treatment and fulfils all of the treatment criteria listed here.	Yes	Yes TA872	28-Feb-23	29-May-23
AXIO1b	Axicabtagene ciloleucel	cell lymphoma (PMBCL) and transformed follicular lymphoma (TFL) to DLBCL in	This application for continuation is being made by and treatment with axicabtagene ciloleucel-modified CAR-T cells will be initiated by a consultant haematologist/medical oncologist specifically trained and accredited in the use of systemic anti-cancer therapy and working in an accredited CAR-T cell treatment centre and who is a member of the National CAR-T clinical Panel for DLBCL, PMBCL and TFL and a member of the treating Trust's DLBCL, PMBCL and TFL and cAR-T cell multidisciplinary teams.  2. The patient has an ECOG performance score of 0 or 1 or 2. Please tick one of the boxes below as to the patient's current ECOG performance status (PS): The ECOG performance status scale is as follows: PS 1 The patient is fully active and able to carry on all pre-disease performance without restriction PS 1 The patient is restricted in physically strenuous activity but is ambulatory and able to carry out work of a light or sedentary nature eg light house work, office work PS 2 The patient is restricted in physically strenuous activity but is ambulatory and able to carry out any work activities and is up and about more than 50% of waking hours PS 3 The patient is capable of only limited selfcare and is confined to bed or chair more than 50% of waking hours PS 4 The patient is completely disabled, cannot carry out any selfcare and is totally confined to bed or chair The patient is completely disabled, cannot carry out any selfcare and is totally confined to bed or chair The patient is completely disabled, cannot carry out any selfcare and is totally confined to bed or chair The patient has required bridging therapy in between leucapheresis and CAR-T cell infusion, please indicate what type(s) of bridging therapy have been required by ticking the most appropriate option below:  - cordioasteroids only or - corticosteroids only or - chemo(immuno)therapy only or - chemo(immuno)therapy or or - chemo(immuno)therapy or and adiotherapy or - chemo(immuno)therapy or and adiotherapy or - chemo(immuno)therapy or adiotherapy or continu	Yes	TA872	28-Feb-23	29-May-23

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ueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with oral azacitidine will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has newly diagnosed acute myeloid leukaemia (AML).				Started
			3. The patient has been treated with standard intensive cytarabine-based induction chemotherapy.				
			4. The patient has either received any consolidation chemotherapy or not. Please mark below whether consolidation chemotherapy was received or not:  - no consolidation chemotherapy was administered - at least one cycle of consolidation chemotherapy was given				
			5. The patient is currently in complete remission (CR) or is in complete remission with incomplete blood count recovery (CRI). Please mark below as to whether the patient is in CR or CRI CR - CR - Cri				
		Oral azacitidine as maintenance therapy in newly diagnosed AML patients in remission following at least induction	6. The patient is not a candidate for, or has chosen not to proceed to, haemopoietic stem cell transplantation (HSCT). Please mark below the reason for not undergoing haemopietic stem cell transplantation: - the patient is not medically lift for HSCT - the patient has chosen not to proceed to HSCT - the patient has chosen not to proceed to HSCT - there is another reason for not proceeding to HSCT			02-Sep-22	
AZA1	Azacitidine	chemotherapy and who are not candidates for, or who choose not to	7. Maintenance therapy with oral azacitidine will be as monotherapy.	No	TA827	05-Oct-22	(Supply
		proceed to, haemopoietic stem cell transplantation where the following	8. Oral azacitidine maintenance therapy will be continued until disease progression up to a <b>maximum of 15% blasts</b> is observed in peripheral blood/bone marrow or until unacceptable toxicity occurs or there is withdrawal of patient consent, whichever is the sooner.				available from 13-Oct-22)
		treatment criteria have been met:	9. The prescribing clinician understands that the usual 300mg once daily 14-day treatment schedule every 28 days for oral azacitidine can be extended to a 21-day treatment schedule every 28 days if a disease relapse with a blast count of 5-15% is observed in the peripheral blood or bone marrow.  Note: oral azacitidine must be discontinued if the blast count exceeds 15% in the peripheral blood or bone marrow.				
			10. The patient is fit for treatment with oral azacitidine maintenance therapy and has an ECOG performance status (PS) of 0-3. Please mark below the ECOG PS status: -PS 0 -PS 1 -PS 1 -PS 2				
		- PS 3					
			1.1. The prescribing climitary transfer and association for a season that or a season that	njectable azacitidine.			
		12. A formal medical review as to whether treatment with oral azactionine should continue will occur at least by the end of the second cycle of treatment.  13. Where a treatment break of more than 10 weeks beyond the expected cycle length is needed, the prescribing clinician will complete a treatment break approval form to restart treatment, including indicating as appropriate if the patient had an extended break because of COVID 19.					
			14. Azacitidine will be otherwise used as set out in its Summary of Product Characteristics (SPC).				
			1. Application made by and first cycle of systemic anti-cancer therapy to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy				
		The first line treatment of low grade	2. Low grade non-Hodgkin's lymphoma		n/a - NHS England		
BEN1	Bendamustine	lymphoma where all the following criteria are met:		Yes	clinical policy	-	08-Jul-18
		are met.	4. To be used within the treating Trust's governance framework, as Bendamustine is not licensed in this indication  Note: Can be used in combination with Rituximab, which is commissioned by NHS England for this indication.	_			
			Note: can be used in combination with intustinal, writer is commissioned by may registed for the induction.  I. Application made by and first cycle of systemic anti-cancer therapy to be prescribed by a consultant specifically trained and accredited in the use of systemic anti-cancer therapy.				
		The first line treatment of mantle cell non-	2. Mantle cell non-Hodgkin's lymphoma				
BEN2	Bendamustine	Hodgkin's lymphoma where all the	3. 1st-line treatment in patients unsuitable for standard treatment	Yes	n/a - NHS England clinical policy	-	08-Jul-18
		following criteria are met:	4. To be used within the treating Trust's governance framework, as Bendamustine is not licensed in this indication		cliffical policy		
			Note: Can be used in combination with Rituximab, which is commissioned by NHS England for this indication.				
			1. Application made by and first cycle of systemic anti-cancer therapy to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy				
			2. Low grade non-Hodgkin's lymphoma 3. Relapsed disease	-			
			5. neinjaeu unbease 4. Unable to receive CHOP-R				
DENIS.		The treatment of relapsed low grade	No. Unable to receive FCR	j	n/a - NHS England		
BEN6	Bendamustine	lymphoma where all the following criteria	6. Unable to receive high dose-therapy	Yes	clinical policy	-	01-Apr-21
		are met:	7. No prior bendamustine	4			
			8. To be used within the treating Trust's governance framework, as Bendamustine is not licensed in this indication	4			
			Note: Can be used in combination with Rituximab, which is commissioned by NHS England for this indication.				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
BEV2	Bevacizumab	The first line treatment of recurrent or metastatic cervical cancer in combination with chemotherapy where all the following criteria are met:	1. An application has been made and the first cycle of systemic anti-cancer therapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The patient has histologically confirmed carcinoma of the cervix  3. The indication will be for 1st line palliative chemotherapy  4. The patient has primary stage IVB, recurrent, or persistent disease not amenable to curative treatment with surgery and/or radiotherapy  5. Bevacizumab will be given with Paclitaxel and either Cisplatin or Carboplatin  6. The patient has an ECOG PS of 0 or 1  7. The patient has had no previous treatment with bevacizumab or other anti-VEGF therapy  8. The patient has no contraindications to the use of bevacizumab  9. Bevacizumab dose to be 15mg/kg every 3 weeks  10. No planned treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve).  *Requests for continuation of treatment after unplanned treatment breaks over this duration should be made via the treatment break approval process  Note: Bevacizumab is ONIV approved for use in combination with combination enthemotherapy and is not approved for use as a single agent maintenance therapy  Note: Bevacizumab should be discontinued for reasons of toxicity or disease progression, whichever occurs first.	Yes	n/a - NHS England clinical policy	-	01-Apr-21
BEV3	Bevacizumab at a dose of 7.5mg/Kg	In combination with 1st line chemotherapy AS INDUCTION TREATMENT for patients with stage III or IV ovarian, fallopian tube or primary peritoneal carcinoma where the following criteria have been met:  Note: there is a separate form BEV9 for the use of bevacizumab at a dose of 15mg/Kg in combination with 1st line chemotherapy AS INDUCTION TREATMENT for advanced ovarian cancer  Note: there is a separate form BEV10 for the use of bevacizumab monotherapy at a dose of 7.5mg/Kg as MAINTENANCE treatment after completion of induction chemotherapy.  Note: there is a separate form OLAP4 for the use of bevacizumab at a dose of 15mg/Kg in combination with olaparib as MaiNTENANCE treatment after completion of induction chemotherapy.	1. This application is being made by and the first cycle of systemic anti-cancer therapy.  2. Bevacizumab at a dose of 7.5mg/kg is to be used in combination with 1st line induction chemotherapy for previously untreated advanced epithelial ovarian, fallopian tube or primary peritoneal cancer.  3. One of the following criteria applies to this patient:  1) FIGO Stage III disease and debulked but residual disease more than 1cm or  1) FIGO Stage III disease and debulked but residual disease more than 1cm or  1) FIGO Stage III disease and unsuitable for debulking surgery or  10 FIGO Stage III disease and unsuitable for debulking surgery or  10 FIGO Stage III disease and unsuitable for debulking surgery or  10 FIGO Stage III disease at presentation and requires neo-adjuvant chemotherapy due to low likelihood of optimal primary surgical cytoreduction  4. Bevacizumab is to be given in combination with carboplatin and paclitaxel chemotherapy.  5. Bevacizumab is to start with:  1) the 1st or 2nd cycle of chemotherapy following primary debulking surgery, or  1) the 1st or 2nd cycle of chemotherapy following interval debulking surgery performed after 3 – 4 cycles of non-bevacizumab-containing neoadjuvant chemotherapy, or  1) the 1st or 2nd cycle of neo-adjuvant chemotherapy  1) the 1st or 2nd cycle of neo-adjuvant chemotherapy  2) the 1st or 2nd cycle of neo-adjuvant chemotherapy  3. As neither this dosage of bevacizumab will be given as part of induction chemotherapy.  4. Bevacizumab is to be given at a dose of 7.5mg/kg every 3 weeks.  7. A maximum of 6 cycles of bevacizumab more its use in the neoadjuvant setting is licensed in ovarian cancer, this use of bevacizumab must be used within the treating Trust's governance framework.  3. As neither this dosage of bevacizumab more its use in the neoadjuvant setting is licensed in ovarian cancer, this use of bevacizumab must be used within the treating Trust's governance framework.  3. As neither this dosage of bevacizumab nor its use in the neoadjuvant setting is licensed in ov	Yes	n/a - NHS England clinical policy	-	01-Apr-21
BEV8	Bevacizumab	The third line treatment of low grade gliomas of childhood where all the following criteria are met:	1. Application made by and first cycle of systemic anti-cancer therapy to be prescribed by a consultant paediatric specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. Progressive low grade glioma  3. No previous treatment with either irinotecan or bevacizumab  4. Irinotecan and bevacizumab to be the 3rd or further line of therapy  5. A maximum of 12 months duration of treatment to be used  6. Consent with the parent/guardian to specifically document the unknown long term toxicity of this combination, particularly on growth and ovarian function  7. To be used within the treating Trust's governance framework, as Bevacizumab and Irinotecan are not licensed in this indication in children  8. In the period immediately prior to the application for irinotecan and bevacizumab, the appropriate specialist MDT has considered the use of proton beam radiotherapy.  NOTE: Bevacizumab is ONLY approved for use in combination with combination chemotherapy and is not approved for use as a single agent maintenance therapy  NOTE: Additional data on long term toxicity must be collected by the paediatric oncology community	Yes			01-Apr-21

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
BEV9	Bevacizumab at a dose of 15mg/Kg	in combination with 1st line chemotherapy AS INDUCTION TREATMENT patients with stage lill or IV ovarian, fallopian tube or primary peritoneal carcinoma where the following criteria have been met.  Note: there is a separate form BEV3 for the use of bevacizumab at a dose of 7.5mg/kg in combination with 1st line chemotherapy AS INDUCTION TREATMENT for advanced ovarian cancer Note: there is a separate form BEV10 for the use of bevacizumab monotherapy at a dose of 7.5mg/kg as MAINTENANCE treatment after completion of induction chemotherapy.  Note: there is a separate form OLAP4 for the use of bevacizumab at a dose of 15mg/kg in combination with olaparib as MAINTENANCE treatment after completion of induction chemotherapy.	1. I confirm that this application is being made by and the first cycle of systemic anti-cancer therapy with bevacizumab in combination with induction chemotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. I confirm that one of the following criteria applies to this patient:  3. I confirm that one of the following criteria applies to this patient:  3. I confirm that one of the following criteria applies to this patient:  3. I confirm that one of the following criteria applies to this patient:  3. I confirm that one of the following criteria applies to this patient:  3. I confirm that one of the following criteria applies to this patient:  3. I confirm that one of the following criteria applies to this patient:  3. I confirm that one of the following criteria applies to this patient:  3. I confirm that bevaciumab is decided with residual disease less than 1 cm or  4. I foll stage II disease and unsuitable for debulking surgery or  5. I confirm that bevaciumab is to be given in combination with carboplatin and paclitaxel chemotherapy.  5. I confirm that bevaciumab is to be given in combination with carboplatin and paclitaxel chemotherapy.  5. I confirm that bevaciumab is to start with:  9. I that or 2nd cycle of chemotherapy following primary debulking surgery, or  10 the 1st or 2nd cycle of chemotherapy following primary debulking surgery, or  10 the 1st or 2nd cycle of chemotherapy following primary debulking surgery, or  10 the 1st or 2nd cycle of chemotherapy following primary debulking surgery, or  11 the 1st or 2nd cycle of chemotherapy following primary debulking surgery, or  12 the 1st or 2nd cycle of chemotherapy following interval debulking surgery, or  13 the 1st or 2nd cycle of chemotherapy following interval debulking surgery, or  14 the 1st or 2nd cycle of hemotherapy following interval debulking surgery or  15 the 1st or 2nd cycle of hemotherapy following interval debulking surgery or  16 the 1st or 2nd cycle of	Yes	n/a - NHS England clinical policy		01-Apr-21
BEV10	Bevacizumab at a dose of 7.5mg/Kg	As MAINTENANCE monotherapy for patients with stage III or IV ovarian, fallopian tube or primary peritoneal carcinoma where the following criteria have been met:  Note: there is a separate form BEV3 for the use of bevacizumab at a dose of 7.5mg/kg in combination with 1st line chemotherapy AS INDUCTION TREATMENT for advanced ovarian cancer  Note: there is a separate form BEV9 for the use of bevacizumab at a dose of 15mg/kg in combination with 1st line chemotherapy AS INDUCTION TREATMENT for advanced ovarian cancer  Note: if an application is being made for the 1st line maintenance combination of olaparia plus bevacizumab, form OLAP4 should be used and will apply to the maintenance use of both drugs	10. I confirm that bevacizumab is to be otherwise used as set out in its Summary of Product Characteristics.  1. I confirm that this application is being made by and the first cycle of systemic anti-cancer therapy with maintenance bevacizumab monotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. I confirm that bevacizumab at a dose of 7.5mg/Kg is to be used as maintenance monotherapy after completion of 1st line induction chemotherapy in combination with bevacizumab 7.5mg/Kg for previously untreated advanced epithelial ovarian, fallopian tube or primary peritoneal cancer.  3. I confirm that this application for maintenance bevacizumab monotherapy continues the use of bevacizumab 7.5mg/Kg previously given in combination with 1st line induction chemotherapy.  4. I confirm that bevacizumab is to be given as monotherapy for a maximum of 18 cycles in all, this figure including the number of cycles given in combination with 1st line induction chemotherapy.  5. I confirm that I understand that this dosage of bevacizumab is not licensed in ovarian cancer, this use of bevacizumab must be used within the treating Trust's governance framework.  Note: This policy relating to the use of maintenance bevacizumab 7.5mg/Kg is NOT for patients with stage I-III disease who have had optimal debulking  7. I confirm that when a treatment break is needed of more than 6 weeks beyond the expected cycle length of 3-weekly treatment, I will complete a treatment break approval form to restart treatment, including as appropriate if the patient had an extended break on account of Covid-19.  8. I confirm that bevacizumab is to be otherwise used as set out in its Summary of Product Characteristics.	Yes	n/a - NHS England clinical policy	-	01-Apr-21

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
BLI1	Blinatumomab	The treatment of relapsed/refractory Philadelphia negative B-precursor acute lymphoblastic leukaemia in ADULT patients	1. An application is being made and the first cycle of systemic anti-cancer therapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient is an adult.  NB. There is a separate Blueteq form to be used for blinatumomab in this indication in children.  3. The patient has relapsed or refractory Philadelphia negative acute lymphoblastic leukaemia (ALL).  4. The patient has been previously treated with intensive combination chemotherapy as initial treatment with or without subsequent salvage therapy.  5. Blinatumomab will only be requested by and administered in either bone marrow transplant centres or in major haematological centres that regularly treat patients with relapsed ALL and who have close and regular ALL multi-disciplinary team meetings and links with bone marrow transplant centres.  6. The patient has an ECOG performance status of 0 - 2.  7. A maximum of 5 cycles of treatment with blinatumomab will be administered.  8. Blinatumomab in this indication is exempt from the NHS England Treatment Break policy.	Yes	TA450	27-Apr-17	26-Sep-17
BLI2	Blinatumomab	The treatment of relapsed/refractory Philadelphia negative B-precursor acute lymphoblastic leukaemia in CHILD patients	9. Blinatumomab will otherwise be used as set out in its Summary of Product Characteristics (SPC).  1. An application is being made and the first cycle of systemic anti-cancer therapy with blinatumomab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient is a child and ONE of the following applies:  OPTION 1 - The patient is post pubescent.  OPTION 2 - The patient is pre pubescent.  Please choose correct option  - Option A  - Option A  - Option B  NB. There is a separate Blueteq form to be used for blinatumomab in this indication in adults.  3. The patient has been previously treated with intensive combination chemotherapy as initial treatment with or without subsequent salvage therapy.  5. The first cycle of blinatumomab will only be requested by, prescribed, and commenced in Principal Treatment Centres (PTCs). Subsequent cycles (including the latter parts of the first 28-day treatment cycle) of blinatumomab may be administered at the PTC or in partnership with enhanced POSCUs under the direction of the PTCs and in agreement with relevant Operational Delivery Networks  6. The use of the blinatumomab has been discussed at a multidisciplinary team (MDT) meeting which must include at least two consultants in the subspecialty with active and credible expertise in the relevant field of whom at least one must be a consultant paediatrician. The MDT should include a paediatric pharmacist and other professional groups appropriate to the disease area.  7. The patient has a Kannofskylarabsy performance score of 60 or more.  8. A maximum of 5 cycles of treatment with blinatumomab will be administered.  9. The use of blinatumomab in this indication is exempt from the NHS England Treatment Break policy.  10. Relevant Trust policy regarding off-label treatments will be followed for children less than 1 year of age, as blinatumomab is not licensed in this age group.	Yes	TA450	27-Apr-17	26-Sep-17

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
виз	Blinatumomab	The treatment of patients in first complete haematological complete remission and with minimal residual disease post 1st line induction chemotherapy in 8- precursor acute lymphoblastic leukaemia in ADULT patients where all the following criteria are met:	1. This application has been made by and the first cycle of systemic anti-cancer therapy with blinatumomab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient is an adult*  *note there is a separate Bluteq form to be used for blinatumomab in this minimal residual disease indication in children.  3. The patient has CD19 positive acute lymphoblastic leukaemia (ALL).  Philadelphia positive ALL (use is on-label) or  Philadelphia negative ALL (use is on-label) or  Philadelphia positive ALL (use is on-label). By ticking this box for use in Philadelphia positive ALL, I confirm that my hospital Trust policy regarding unlicensed treatments is being followed as blinatumomab is not licensed in Philadelphia positive ALL.  4. The patient has been previously treated with intensive 1st line combination chemotherapy as initial induction treatment.  5. The patient's bone marrow has been shown to have a minimal residual disease level of ≥ 0.01% (≥10-4) leukaemic cells confirmed in a validated assay.  Note: a patient who has minimal residual disease (MRD) negativity defined as being less than 0.01% is potentially eligible for blinatumomab as part of consolidation therapy via form BLIS.  7. Blinatumomab will only be requested by and administered in either bone marrow transplant centres or in major haematological centres that regularly treat patients with MRD positive ALL and who have close and regular ALL multidisciplinary team meetings and links with bone marrow transplant centres.  8. The patient has an £COS performance status of 0-2.  9. A maximum of 4 cycles of blinatumomab will be administered to this patient.  10. Blinatumomab will be used as set out in its Summary of Product Characteristics (SPC).	No	TA589	24-Jul-19	22-Oct-19
BLI4	Blinatumomab	induction chemotherapy in B-precursor acute lymphoblastic leukaemia in CHILD patients where all the following criteria have been met:	1. This application has been made by and the first cycle of systemic anti-cancer therapy with blinatumomab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient is a child⁴ and please mark as to whether pre- or post-pubescent:  1. is post-pubescent or  2. The patient has CD19 sostitive acute lymphoblastic leukaemia (ALL).  3. The patient has CD19 sostitive acute lymphoblastic leukaemia (ALL).  Philadelphia negative ALL or  Philadelphia negative ALL or  Philadelphia positive ALL  4. The patient has been previously treated with 1st line intensive combination chemotherapy as initial induction treatment.  5. The patient is in complete haematological remission of ALL.  6. The patient's bone marrow has been shown to have minimal residual disease level of ≥ 0.01% (210-⁴) confirmed in a validated assay.  Note: a patient who has minimal residual disease (MRD) negativity defined as being less than 0.01% is potentially eligible for blinatumomab as part of consolidation therapy via form BLI6.  7. The first cycle of blinatumomab will only be requested by, prescribed, and commenced in Principal Treatment Centres (PTCs). Subsequent cycles (including the latter parts of the first 28-day treatment cycle) of blinatumomab may be administered at the PTC or in partnership with enhanced POSCUs under the direction of the PTCs and in agreement with relevant Operational Delivery Networks.  8. The patient has a Karnofisky/Lancksy performance score of 60 or more.  9. A maximum of 4 cycles of treatment with blinatumomab will be administered.  10. Blinatumomab has been discussly performance score of 60 or more.  9. A maximum of 4 cycles of treatment with blinatumomab will be administered.  10. Blinatumomab has been discussed at a multi-disciplinary team (MDT) meeting which must include at least two consultants in the subspecialty with active and credible expertise in the relevant field of whom at least one must be a consultant paediatrician. The MDT sh	No	TA589	24-Jul-19	22-Oct-19

ilueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with blinatumomab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			2. The patient is an adult.	-			
			3. The patient has Philadelphia chromosome negative CD19 positive B-cell precursor acute lymphoblastic leukaemia (ALL).				
			4. The patient has been previously treated with intensive 1st line induction and intensification combination chemotherapy.				
			5. The patient is in a morphological complete remission of ALL.	-			
			6. The prescribing clinician understands that this NICE recommendation for blinatumomab uses the E1910 trial definition of minimal residual disease negativity as the bone marrow exhibiting <0.01% (<10.4) leukaemic cells confirmed in a validated assay and the prescibing clinician confirms that this patient's level of minimal residual disease fulfils this definition. For those patients in whom an assay sensitivity or QR of 10.4 is not reached but sufficient to report minimal residual disease negativity to the maximum sensitivity of the available assay, blinatumomab will also be permitted.				
			Note: the company's case for the clinical and cost effectiveness of blinatumomab in this indication was based on the evidence base of the E1910 trial which in the key randomisation only included patients who had MRD negativity defined as being <0.01%.				
		The second secon	indication which can be accessed via form BLI3.				
BLI5	Blinatumomab	chemotherapy for Philadelphia	7. Blinatumomab will only be requested by and administered in either bone marrow transplant centres or in major haematological centres that regularly treat patients with MRD negative ALL and who have close and regular ALL multi-disciplinary team meetings and links with bone marrow transplant centres.	No	TA1049	26-Mar-25	24-Jun-25
		chromosome negative B-cell precursor acute lymphoblastic leukaemiawhere all	8. The patient has an ECOG performance status of 0-2.	1		1	
		the following criteria are met:	9. The treatment intent for this patient is to be potentially treated with a maximum of 4 cycles of blinatumomab whether given in cycles 1, 2, 6 and 8 of consolidation treatment with chemotherapy planned to be given in cycles 3, 4, 5 and 7 of an 8 cycle consolidation treatment program or blinatumomab given in cycles 1, 2, 6 and 7 and chemotherapy in cycles 3, 4 and 5 of a 7 cycle consolidation treatment program or blinatumomab as sequenced with chemotherapy in other approved UK ALL Research Network consolidation treatment protocols.	-			
			Note: NHS England understands that patients in the E1910 trial could proceed to allogeneic transplantation after completing at least cycles 1 and 2 of the above potential program of consolidation therapy.				
			10. The patient has not yet commenced any consolidation therapy i.e. the patient has just finished the sequence of induction and intensification therapies.				
			Note: the company's case for the clinical and cost effectiveness of blinatumomab in this indication was based on the evidence base of the E1910 trial which only included patients who had not started any consolidation therapy.				
			11. Blinatumomab will be administered as monotherapy in accordance with treatment criterion 9 above.	1			
			Nato: intersheed showsthereous and nonconsists turnsing kinese inhibitory (for patients with ADI shore mutations) was to continued as also and during any billiontum mark or also				
			Note: intrathecal chemotherapy and appropriate tyrosine kinase inhibitors (for patients with ABL-class mutations) may be continued as planned during any blinatumomab cycles.  12. The prescribing clinician understands that given the scheduling timetable of a potential maximum of 4 cycles of blinatumomab given interspersed with cycles of chemotherapy, this indication is exempted from NHS England's				
			treatment break policy.				
			13. Blinatumomab will otherwise be used as set out in its Summary of Product Characteristics (SPC).				
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with blinatumomab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			2. The patient is a post pubescent child.	1			
			3. The patient has Philadelphia chromosome negative CD19 positive B-cell precursor acute lymphoblastic leukaemia (ALL).	-			
			4. The patient has been previously treated with intensive 1st line induction and any indicated cytoreductive combination chemotherapy.				
			5. The patient is in a morphological complete remission of ALL.	1			
			6. The prescribing clinician understands that this NICE recommendation for blinatumomab uses the E1910 trial definition of minimal residual disease negativity as the bone marrow exhibiting <0.01% (<10.4) leukaemic cells confirmed in a validated assay and the prescibing clinician confirms that this patient's level of minimal residual disease fulfils this definition. For those patients in whom an assay sensitivity or QR of 10.4 is not reached but sufficient to report minimal residual disease negativity to the maximum sensitivity of the available assay, blinatumomab will also be permitted.	-			
		The treatment of POST PUBESCENT	Note: the company's case for the clinical and cost effectiveness of blinatumomab in this indication was based on the evidence base of the E1910 trial which in the key randomisation only included patients who had MRD negativity defined as being <0.01%.				
		CHILDREN in first morphological complete	Note: a level of minimal residual disease (MRD) of >=0.01% means that blinatumomab is not recommended by NICE in this indication and is not funded by NHS England. Blinatumomab is however potentially funded in a MRD positive indication which can be accessed via form BLI4.				
BLI6	Blinatumomab	and any indicated intensification chemotherapy for Philadelphia chromosome negative B-cell precursor	7. Blinatumomab will only be requested by, prescribed, and initially administered in, principal treatment centres (PTCs) who have close and regular ALL multi-disciplinary team meetings and links with bone marrow transplant centres. Subsequent cycles of blinatumomab (including the latter part of the first 28-day treatment cycle) may be administered at PTCs or in close partnership with enhanced POSCUs under the direction of PTCs and in agreement with relevant Operational Delivery Networks.	No	TA1049	26-Mar-25	24-Jun-25
		acute lymphoblastic leukaemia where all	8. The patient has a Karnofsky/Lansky performance score of at least 60.				
		the following criteria have been met:	9. The treatment intent for this patient is to be potentially treated with a maximum of 4 cycles of blinatumomab as sequenced with chemotherapy in accordance with UK nationally approved CCLG protocols/guidelines.  Note: NHS England understands that patients in the E1910 trial could proceed to allogeneic transplantation after completing at least cycles 1 and 2 of blinatumomab consolidation therapy.				
			10. The patient has not yet commenced any consolidation therapy i.e. the patient has just finished the sequence of induction and any indicated cytoreductive therapies.  Note: the company's case for the clinical and cost effectiveness of blinatumomab in this indication was based on the evidence base of the E1910 trial which only included patients who had not started any consolidation therapy.				
			11. Blinatumomab will be administered as systemic monotherapy in accordance with treatment criterion 9 above.	_			
			Note: intrathecal chemotherapy, , and appropriate tyrosine kinase inhibitors, may continue as planned during blinatumomab cycles.				
			12. The prescribing clinician understands that given the scheduling timetable of a potential maximum of 4 cycles of blinatumomab given interspersed with cycles of chemotherapy, this indication is exempted from the NHS England's treatment break policy.				
			13. Trust policy regarding unlicensed treatments has been followed as blinatumomab is not licensed in this indication in post pubescent children.				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
BOS1	Bosutinib	Bosutinib for previously treated chronic myeloid leukaemia	1.1 confirm that an application has been made and the first cycle of systemic anti-cancer therapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2.1 confirm the patient has chronic, accelerated or blast phase Philadelphia chromosome positive chronic myeloid leukaemia.  3.1 confirm the patient has had previous treatment with 1 or more tyrosine kinase inhibitor.  4.1 confirm that treatment is not appropriate with either imatinib, nilotinib or dasatinib.  5.1 confirm the patient will receive the licensed dose and frequency of bosutinib	Yes	TA401	24-Aug-16	22-Nov-16
BRE3 (formerly BRE2)	Brentuximab	Treatment of brentuximab-naïve relapsed/refractory Hodgkin lymphoma following autologous stem cell transplant in ADULT patiests where the following criteria are met:	1. An application has been made by and the first cycle of systemic anti-cancer therapy is to be prescribed by a consultant specifically trained and accredited in the use of systemic anti-cancer therapy  2. The patient is an adult.  NB. There is a separate Blueteq form to be used for brentuximab in this indication in children.  3. The patient has relapsed or refractory CD30+ Hodgkin lymphoma.  4. The patient has relapsed Hodgkin lymphoma after autologous stem cell transplant.  5. The patient has relapsed Hodgkin lymphoma after autologous stem cell transplant.  5. The patient has never received brentuximab unless having previously responded to brentuximab when treated with 1st line BV-AVD.  6. Treatment with brentuximab will be discontinued after 4 cycles if CT or PET-CT scans to assess response demonstrate a response status of less than a partial or a complete response  7. No planned treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve).  *Requests for continuation of treatment after unplanned treatment breaks over this duration should be made via the treatment break approval process  8. No re-use of brentuximab will be used outside this indication unless previous partial/complete response to brentuximab and brentuximab is being used as a bridge to allogeneic stem cell transplant or donor lymphocyte infusion*  *note there is a separate blueteq form for such re-use of brentuximab  9. A maximum of 16 cycles of brentuximab will be administered to the patient  Note: administration of a full 6 cycles of 1st line use of 8b y lbus AVD (12 doses of brentuximab at 1.2 mg/kg) counts as 8 cycles of brentuximab monotherapy at 1.8mg/kg.  10. Brentuximab will otherwise be used as set out in its Summary of Product Characteristics (SPC).	Yes	TA524 (formerly TA446)	13-Jun-18	26-Sep-17
BRE4 (formerly BRE2)	Brentuximab	Treatment of brentuximab-naïve relapsed/refractory Hodgkin lymphoma following autologous stem cell transplant in CHILD patients where the following criteria are met:	1. An application has been made by and the first cycle of systemic anti-cancer therapy is to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The patient has relapsed or refractory CD30+ Hodgkin lymphoma.  3. The patient has relapsed Hodgkin lymphoma after autologous stem cell transplant  4. The patient has never received brentuximab  5. Treatment with brentuximab will be discontinued after 4 cycles if CT or PET-CT scans to assess response demonstrate a response status of less than a partial or a complete response  6. The patient is a child* and is either post pubescent or is pre pubescent and will receive brentuximab dosage described in the phase 2 part of the brentuximab trial protocol C25002  http://www.clinicaltrials.gov/ct2/show/NCT014920887term=C25002&rank=1 and reported on http://www.bloodjournal.org/content/122/21/4378  *note there is a separate Bluteq form to be used for brentuximab in this indication in adults.  7. The use of the brentuximab has been discussed at a multi disciplinary team (MDT) meeting which must include at least two consultants in the subspecialty with active and credible expertise in the relevant field of whom at least one must be a consultant paediatrician. The MDT should include a paediatric pharmacist and other professional groups appropriate to the disease area.  8. No planned treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve).  *Requests for continuation of treatment after unplanned treatment breaks over this duration should be made via the treatment break approval process  9. No re-use of brentuximab will be used outside this indication unless previous partial/complete response to brentuximab and brentuximab is being used as a bridge to allogeneic stem cell transplant or donor lymphocyte infusion*  **note there is a separate blueteq form for such re-use of brentuximab  10. A max	Yes	TA524 (formerly TA446)	13-Jun-18	26-Sep-17

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. Application made by and first cycle of systemic anti-cancer therapy to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy				
			The patient is an adult*     *note there is a separate blueteq form to be used for brentuximab in this indication in children				
			3. The patient has relapsed or refractory CD30+ Hodgkin lymphoma.				
			4. The patient has relapsed Hodgkin lymphoma after at least 2 prior systemic therapies when either autologous stem cell transplant or further multi-agent chemotherapy is not a treatment option.				
			5. The patient has had no previous stem cell transplant				
		Treatment of brentuximab-naïve	6. The The patient has never received brentuximab unless having previously responded to brentuximab when treated with 1st line BV-AVD.				
		relapsed/refractory Hodgkin lymphoma	- No prior treatment with brentuximab				
BRE5		following at least 2 prior therapies when	- Prior therapy brentuximab within 1st line BV-AVD				
(formerly BRE2)	Brentuximab	autologous stem cell transplant or multi-	7. Treatment with brentuximab will be discontinued after 4 cycles if CT or PET-CT scans to assess response demonstrate a response status of less than a partial or a complete response	Yes	TA524	13-Jun-18	11-Sep-18
		agent chemotherapy is not a treatment option in ADULT patients where the	8. I confirm that no more than 16 cycles of brentuximab may be administered per patient				
		following criteria are met:	Note: administration of a full 6 cycles of 1st line use of BV plus AVD (12 doses of brentuximab at 1.2 mg/kg) counts as 8 cycles of brentuximab monotherapy at 1.8 mg/kg.				
		3	9. No planned treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve).	-			
			Requests for continuation of treatment after unplanned treatment breaks over this duration should be made via the treatment break approval process				
			10. No re-use of brentuximab will be used outside this indication unless previous partial/complete response to brentuximab and brentuximab is being used as a bridge to allogeneic stem cell transplant or donor lymphocyte infusion*				
			*note there is a separate blueteq form for such re-use of brentuximab				
			11. Brentuximab will otherwise be used as set out in its Summary of Product Characteristics (SPC).	-			
			1. Application made by and first cycle of systemic anti-cancer therapy to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy				
			2. The patient is a child* and is either post pubescent or is pre pubescent and will receive brentusimab dosage described in the phase 2 part of the brentusimab trial protocol C25002				
			http://www.clinicaltrials.gov/ct2/show/NCT01492088?term=C25002&rank=1 and reported on http://www.bloodjournal.org/content/122/21/4378				
			*note there is a separate Bluteq form to be used for brentuximab in this indication in adults.				
			3. The patient has relapsed or refractory CD30+ Hodgkin lymphoma.	-	TAS24		
			4. The patient has relapsed to depkin lymphoma after at least 2 prior systemic therapies when either autologous stem cell transplant or further multi-agent chemotherapy is not a treatment option.	1			
			5. The patient has had no previous stem cell transplant	-			
		Treatment of brentuximab-naïve	5. The patient has never received brentusmab				
		relapsed/refractory Hodgkin lymphoma	7. Treatment with brentuximab will be discontinued after 4 cycles if CT or PET-CT scans to assess response demonstrate a response status of less than a partial or a complete response				
BRE6	Brentuximab	following at least 2 prior therapies when autologous stem cell transplant or multi-	8. I confirm that no more than 16 cycles of brentuximab may be administered per patient	Yes	TAS24 13-Jun-18	13-Jun-18	11-Sep-18
(formerly BRE2)		agent chemotherapy is not a treatment	9. The use of the brentuximab has been discussed at a multi disciplinary team (MDT) meeting which must include at least two consultants in the subspecialty with active and credible expertise in the relevant field of whom at least one	-			
		option in CHILD patients where the	must be a consultant paediatrician. The MDT should include a paediatric pharmacist and other professional groups appropriate to the disease area.				
		following criteria are met:	10. No planned treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve).	-			
			*Requests for continuation of treatment after unplanned treatment breaks over this duration should be made via the treatment break approval process				
			11. No re-use of brentuximab will be used outside this indication unless previous partial/complete response to brentuximab and brentuximab is being used as a bridge to allogeneic stem cell transplant or donor lymphocyte infusion*	1			
			*note there is a separate blueteq form for such re-use of brentuximab				
			12. Trust policy regarding unlicensed treatments has been followed as brentuximab is not licensed in this indication in children.	4			
			13. Brentuximab will otherwise be used as set out in its Summary of Product Characteristics (SPC).				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
BRE7	Brentuximab	Re-use of brentuximab in relapsed/refractory Hodgkin lymphoma in ADULT patients:	1. An application has been made by and the first cycle of systemic anti-cancer therapy is to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The patient has relapsed or refractory CD30+ Hodgkin lymphoma.  3. The patient has relapsed Hodgkin lymphoma after autologous stem cell transplant  4. Previous use of brentuximab achieved a partial/complete response to brentuximab  5. Brentuximab is being used as a bridge to allogeneic stem cell transplantation or donor lymphocyte infusion  6. Treatment with brentuximab will be discontinued after 4 cycles if CT or PET-CT scans to assess response demonstrate a response status of less than a partial or a complete response  7. The patient is an adult*  **once there is a separate blueteq form to be used for brentuximab in this indication in children  8. No planned treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve).  *Requests for continuation of treatment after unplanned treatment breaks over this duration should be made via the treatment break approval process  9. A maximum of 16 cycles of brentuximab will be administered to the patient when combining this reuse and previous cycles of brentuximab monotherapy at 1.8mg/kg.  10. Brentuximab will otherwise be used as set out in its Summary of Product Characteristics (SPC).	Yes	TA524 (formerly TA446)	13-Jun-18	26-Sep-17
BRE8	Brentuximab	Re-use of brentuximab in relapsed/refractory Hodgkin lymphoma in CHILD patients:	1. An application has been made by and the first cycle of systemic anti-cancer therapy is to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The patient has relapsed or refractory CD30+ Hodgkin lymphoma.  3. The patient has relapsed Hodgkin lymphoma after autologous stem cell transplant  4. Previous use of brentuximab achieved a partial/complete response to brentuximab  5. Brentuximab is being used as a bridge to allogeneic stem cell transplantation or donor lymphocyte infusion  6. Treatment with brentuximab will be discontinued after 4 cycles if CT or PET-CT scans to assess response demonstrate a response status of less than a partial or a complete response  7. The patient is a child* and is either post pubescent or is pre pubescent and will receive brentuximab dosage described in the phase 2 part of the brentuximab trial protocol C25002 http://www.clinicaltrials.gov/ct2/show/NCT014920887term=C250028rank=1 and reported on http://www.bloodjournal.org/content/122/21/4378  **note there is a separate Bluteq form to be used for brentuximab in this indication in adults.  8. The use of the brentuximab has been discussed at a multi disciplinary team (MDT) meeting which must include at least two consultants in the subspecialty with active and credible expertise in the relevant field of whom at least one must be a consultant paediatrician. The MDT should include a paediatric pharmacist and other professional groups appropriate to the disease area.  9. No planned treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve).  *Requests for continuation of treatment after unplanned treatment breaks over this duration should be made via the treatment break approval process  10. A maximum of 16 cycles of brentuximab will be administered to the patient when combining this reuse and previous cycles of brentuximab  11. Trust policy regardi	Yes	TA524 (formerly TA446)	13-Jun-18	26-Sep-17

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is being made by and first cycle of systemic anti-cancer therapy with brentuximab vedotin will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			2. The patient has relapsed or refractory systemic anaplastic large cell lymphoma (sALCL) after front line chemotherapy.  NB. Brentuximab is not available for primary cutaneous anaplastic large cell lymphoma unless it has transformed into systemic anaplastic large cell lymphoma.				
			3. The patient has a proven histological diagnosis of CD30+ve systemic anaplastic large cell lymphoma.				
BRE9 _		The treatment of relapsed or refractory	4. Either the patient has never previously been treated with brentuximab vedotin or was previously treated with brentuximab vedotin in combination with cyclophosphamide, doxorubicin and prednisone and did not have refractory disease to this therapy.  Please mark which of these 2 clinical scenarios applies to this patient:  - No prior treatment with prentuximab vedotin  - Received prior treatment with brentuximab vedotin in combination with cyclophosphamide, doxorubicin and prednisone and did not have refractory disease to this therapy				
(formerly BRE1)	Brentuximab	systemic anaplastic large cell lymphoma in ADULT patients, where the following	5. Brentwimab is to be used as single-agent therapy.	Yes	TA478	04-Oct-17	02-Jan-18
		criteria have been met:	6. The patient has an ECOG performance status of 0 or 1 or 2.				
		7. Treatment with brentuximab is to be discontinued after 4 cycles if the CT or PET-CT scans to assess response demonstrate a response status of less than a partial or a complete response.					
			8. A maximum of 16 cycles of brentuximab vedotin may be administered per patient (this total of 16 cycles includes any previous treatment with brentuximab vedotin as part of prior therapy).				
			9. A formal medical review as to how the brentuximab vedotin is being tolerated and whether treatment with brentuximab vedotin should continue or not will be scheduled to occur at least by the end of the first 6 weeks of treatment.				
			10. If a treatment break of more than 6 weeks beyond the expected 3 week cycle length occurs, I will complete a treatment break approval form to restart treatment.				
		11. Brentuximab will be otherwise used as set out in its Summary of Product Characteristics (SPC).					
			1. An application has been made and the first cycle of systemic anti-cancer therapy is to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy				
			2. The patient has relapsed or refractory <b>systemic</b> anaplastic large cell lymphoma after front line chemotherapy  Note: Brentuximab is not available for 1° cutaneous anaplastic large cell lymphoma unless it has transformed into <b>systemic</b> anaplastic large cell lymphoma				
			3. Histologically confirmed CD30 positive disease				
			4. The patient has never previously received brentuximab unless previously enrolled in the NCRI-adopted clinical trial called ECHELON-2				
			5. Brentuximab is to be used as single-agent therapy				
			6. The patient has an ECOG performance status of 0-1				
BRE10	Brentuximab	The treatment of relapsed or refractory systemic anaplastic large cell lymphoma in	7. The patient is a child <sup>4</sup> and either post pubescent or is pre pubescent and will receive brentuximab vedotin dosage as described in phase 2 of the trial protocol C25002 http://www.clinicaltrials.gov/ctz/show/NCT014920887rem=C250028rank=1 and reported on http://www.bloodjournal.org/content/122/21/4378  Note: there is a separate Blueteq form to be used for brentuximab vedotin in this indication in addition.	Yes	TA478	04-Oct-17	02-Jan-18
(formerly BRE1)	Dielitaxillab	CHILD patients, where the following criteria have been met:	8. The use of brentuximab in this setting and in this patient has been discussed at a multi-disciplinary team (MDT) meeting which must include at least 2 consultants in the subspecialty with active and credible expertise in the relevant field of whom one must be a consultant paediatrician. The MDT should include a paediatric pharmacist and other professional groups appropriate to the disease area		18470	0.4 0.0 27	02 3411 10
			9. Treatment with brentuximab to be discontinued after 4 cycles if CT or PET-CT scans to assess response demonstrate a response status of less than a partial or a complete response				
			10. No treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve)*				
			*Note: Requests for continuation of treatment after unplanned treatment breaks over this duration should be made via the treatment break approval process				
			11. Brentuximab vedotin will only be requested by and administered in principal treatment centres				
			12.Trust policy regarding unlicensed treatments has been followed as brentuximab vedotin is not licensed in this indication in children				
			13. A maximum of 16 cycles of brentuximab may be administered per patient				
			14. Brentuximab will be otherwise used as set out in its Summary of Product Characteristics			1	L

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
BRE11	Brentuximab vedotin	lymphoma following at least 1 prior	1. This application has been made by and the first cycle of systemic anti-cancer therapy.  2. The patient has relapsed or refractory CD30+ cutaneous T cell lymphoma, the type of which is one of the following: advanced stage IIB-IVB mycosis fungoides or primary cutaneous anaplastic large cell lymphoma or Sezary syndrome Please mark in the tick box below which of these 3 types of cutaneous T cell lymphoma applies to this patient:  - stage IIB-IVB mycosis fungoides or - primary cutaneous anaplastic large cell lymphoma or - sezary syndrome  Note: Takeda restricted its submission to NICE for the consideration of the clinical and cost effectiveness of brentuximab vedotin in only these 3 subtypes of cutaneous T cell lymphoma (CTCL) and NICE has optimised its recommendations in CTC1 accordingly. Brentuximab vedotin is therefore not approved for use in patients with other types of cutaneous lymphoma such as lymphomatoid papulosis, subcutaneous panniculitis-like T cell NHL and primary cutaneous peripheral T cell lymphoma.  3. The patient has been treated with at least 1 prior systemic therapy for his/her CTCL.  4. The patient has never previously received treatment with brentuximab vedotin unless it has been given as part of any compassionate use scheme and the patient meets all the other criteria set out here including the maximum treatment duration of 16 cycles as set out in brentuximab vedotin values it has been given as part of any compassionate use scheme and the patient meets all the other criteria set out here including the maximum treatment duration of 16 cycles of brentuximab vedotin will be administered to this patient.  5. No more than 16 cycles of brentuximab vedotin will be administered to this patient.  6. The patient has an ECOS performance status of 0 or 1 or 2.  7. No planned treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve).  *Requests for continuation of treatment after unplann	No No	TA577	24-Apr-19	23-Jul-19
BRE12	Brentuximab vedotin	The treatment of CD30+ cutaneous T cell lymphoma following at least 1 prior systemic therapy in CHILD patients where the following criteria are metre.  Note: there is a separate Blueteq form for the use of brentuximab vedotin in adults with cutaneous T cell lymphoma	1. This application has been made by and the first cycle of systemic anti-cancer therapy.  2. The patient is a child* and please mark as to whether the child is pre- or post-pubescent:  1. Spost-pubescent or  1. Spost-pubescent and will receive brentuximab vedotin at the paediatric dosage described in the brentuximab vedotin literature in Hodgkin lymphoma.  1. The patient has relapsed or refractory CD30+ cutaneous T cell lymphoma which is advanced stage IIB-IVB mycosis fungoides or primary cutaneous anaplastic large cell lymphoma or Sezary syndrome.  2. The patient has relapsed or refractory CD30+ cutaneous T cell lymphoma applies to this patient:  3. The patient has relapsed or refractory CD30+ cutaneous T cell lymphoma applies to this patient:  3. The patient has relapsed or lefractory CD30+ cutaneous T cell lymphoma applies to this patient:  3. The patient has relapsed or refractory CD30+ cutaneous T cell lymphoma applies to this patient:  3. The patient has relapsed or refractory CD30+ cutaneous T cell lymphoma applies to this patient:  3. The patient has relapsed or refractory CD30+ cutaneous T cell lymphoma applies to this patient:  3. The patient has relapsed or refractory CD30+ cutaneous T cell lymphoma applies to this patient:  3. The patient has relapsed or refractory CD30+ cutaneous T cell lymphoma applies to this patient:  5. Sezary syndrome.  Note: Takedar restricted its submission to NICE for the consideration of the clinical and cost effectiveness of only these 3 subtypes of cutaneous T cell lymphoma (CTCL) and NICE has restricted its recommendations in CTCL accordingly.  Brentuxinably evedotin is therefore not approved for use in patients with other types of cutaneous lymphoma such as lymphomatoid papulosis, subcutaneous paniculitis-like T cell NHL and primary cutaneous peripheral T cell lymphoma.  4. The patient has never previously received brentuximab vedotin unless it has been given as part of a compassionate access scheme and the patient meets all the criteria set out here including the max	No	TA577	24-Apr-19	23-Jul-19

lueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with brentuximab vedotin in combination with chemotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			2. The patient has a proven histological diagnosis of CD30+ve systemic anaplastic large cell lymphoma (sALCL).				
			3. The patient is previously untreated for systemic anaplastic large cell lymphoma.				
	Brentuximab vedotin	For previously untreated systemic	4. The patient has not received prior treatment with brentusimab vedotin.				
BRE13	in combination with	anaplastic large cell lymphoma (sALCL) in	5. The patient will be treated with brentuximab vedotin in combination with cyclophosphamide, doxorubicin and prednisone.	No	TAC41	12-Aug-20	10-Nov-20
BKE13	cyclophosphamide, doxorubicin and	an ADULT patient where the following	6. The patient will be treated with a maximum of 6 or 8 cycles of chemotherapy, 6 cycles being the usual maximum.	, NO	1Ab41		10-NOV-20
	prednisone	criteria have been met:	7. The patient has an ECOG performance status of 0 or 1 or 2.				
			8. A formal medical review as to how the combination of brentuximab vedotin and chemotherapy is being tolerated and whether treatment with the combination of brentuximab vedotin and chemotherapy should continue or not will be scheduled to occur at least by the end of the first 6 weeks of treatment.				
			9. When a treatment break of more than 6 weeks beyond the expected 3 week cycle length occurs, I will complete a treatment break approval form to restart treatment.				
			10. Brentuximab vedotin will otherwise be used as set out in its Summary of Product Characteristics (SPC)				
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with brentuximab vedotin in combination with chemotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			2. The patient has a proven histological diagnosis of CD30+ve systemic anaplastic large cell lymphoma (sALCL).				
			3. The patient is previously untreated for systemic anaplastic large cell lymphoma.		TAG41		
			4. The patient is a child* and the prescribing clinician understands that the Summary of Product Characteristics (SPC) states 'The safety and efficacy in children less than 18 years have not yet been established.'				
BRE14	Brentuximab vedotin in combination with	For previously untreated systemic anaplastic large cell lymphoma (sALCL) in CHILD patients where the following	5. The patient has not received prior treatment with brentusimab vedotin or previous cytotoxic chemotherapy*.  *Note: patients who present with rapidly progressing disease may receive a single course of chemotherapy, as an emergency treatment given before final diagnosis is established.	No	TA641	12-Aug-20	03-Feb-23
	chemotherapy	criteria are met:	6. the patient will be treated with brentuximab vedotin in combination with chemotherapy using the brentuximab vedotin dose (1.8mg/kg) and chemotherapy schedule described in the reference below and I understand that that the trial excluded patients less than 10kg so brentuximab must only be given to patients who weigh 10kg or more.  **Tower Renly AF, Izm MS, Gross TG, Saguillig I, Brokasuskas D et al Brentuximab vedotin in combination with chemotherapy for pediatric patients with ALK1 ALCL: results of COG trial ANHL12P1: Blood 1 July 2021 Volume 137, Number 26,p3595-3603'				
			7. The use of the brentuximab vedotin has been discussed at a multi-disciplinary team (MDT) meeting which must include at least two consultants in the subspecialty with active and credible expertise in the relevant field of whom at least one must be a consultant paediatrician. The MDT should include a paediatric pharmacist and other professional groups appropriate to the disease area.				
			8. The patient has an ECOG (or equivalent Karnofsky/Lansky Scale) performance status of 0 - 2.	1			
			9. The patient does not have disease isolated to the skin, stage I disease, or central nervous system involvement.	]			
			10. Trust policy regarding unlicensed treatments is being followed.	1			
			11. When a treatment break of more than 6 weeks beyond the expected 3 week cycle length occurs, a treatment break approval form will be completed to restart treatment.  *Requests for continuation of treatment after unplanned treatment breaks over this duration should be made via the treatment break approval process				
			12. Brentuximab vedotin will otherwise be used as set out in its Summary of Product Characteristics (SPC).				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
BRE15	Brentuximab vedotin in combination with doxorubicin, vinblastine and dacarbazine	For treating adult patients with previously untreated stage III or IV Hodgkin lymphoma where the following criteria have been met:	1. This application is being made by and the first cycle of brentuximab in combination with doxorubicin, vinblastine and dacarbazine will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient is an adult.  3. The patient has previously untreated CD30 positive Hodgkin lymphoma.  4. The patient has stage III or IV Hodgkin lymphoma.  Please mark below which stage applies to this patient: - stage III disease or - stage IV disease  Note: the use of brentuximab plus chemotherapy is not commissioned in stage I or II Hodgkin lymphoma.  5. Brentuximab will be given in combination with doxorubicin, vinblastine and dacarbazine (AVD).  6. A maximum of 6 x 28 day cycles of brentuximab plus AVD will be administered to this patient.  Note: there is no PET-adapted approach to treatment escalation or de-escalation with this brentuximab-AVD combination.  7. The prescribing clinician is aware that the scheduled brentuximab dose per day 1 and day 15 administrations is 1.2mg/kg (ie not the dose used when brentuximab is given as monotherapy).  8. The prescribing clinician is aware that the brentuximab SPC recommends that primary prophylaxis with GCSF should begin with the first dose of brentuximab-AVD.  9. The patient has an ECOG performance status of 0 or 1 or 2.  10. The prescribing clinician is aware that then a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form.	No	TA1059	07-May-25	05-Aug-25
			11. Brentuximab will be otherwise used as set out in its Summary of Product Characteristics (SPC).				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
BRII	Brigatinib	Brigatinib for anaplastic lymphoma kinase- positive advanced non-small-cell lung cancer previously treated with crizotinib where all the following criteria have been met:	1. This application is made by and the first cycle of systemic anti-cancer therapy with brigatinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has histological or cytological evidence of NSCLC that carries an anaplastic lymphoma kinase (ALK) rearrangement based on a validated test <u>OR</u> there is documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC AND there is an informative circulating free DNA test result confirming the presence of an activating anaplastic lymphoma kinase (ALK) rearrangement.  Please mark below on which basis the diagnosis of ALK positive NSCLC has been made in this patient:  - Histological or cytological evidence Documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC and there is an informative circulating free DNA test result confirming the presence of an activating anaplastic lymphoma kinase (ALK) rearrangement.  3. The only TKI treatment that the patient has progressed on is 1st line crizotinib or 2nd line crizotinib after 1st line chemotherapy and that the patient has not been treated with either 1st line alectinib or 1st line ceritinib.  Second line brigatinib is only licensed, NICE-approved and funded in patients who have been treated with and progressed on crizotinib as their sole TKI treatment.  4. The patient has not been treated with 2nd line ceritinib after 1st line crizotinib unless the ceritinib had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease nonzeaszalom.  5. The patient has not been previously treated with brigatinib unless brigatinib has been received as part of any compassionate use scheme and the patient meets all the other criteria set out here.  6. Brigatinib will be used only as monotherapy.  7. The patient has no brain metastases or, if the pa	No -	TA571	20-Mar-19	18-Jun-19
BR/2_v1.3	Brigatinib monotherapy	For anaplastic lymphoma kinase-positive advanced non-small cell lung cancer previously untreated with an ALK inhibitor where the following criteria have been met:	1. This application for brigatinib is being made by and the first cycle of systemic anti-cancer therapy.  2. The patient has locally advanced or metastatic non-small cell lung cancer.  3. The patient has locally advanced or metastatic non-small cell lung cancer.  3. The patient has locally advanced or metastatic non-small cell lung cancer.  3. The patient has bistological or cyclological evidence of NSCLC that carries an anaplastic lymphoma kinase (ALK) rearrangement based on a validated test <b>QR</b> there is documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC <b>AND</b> there is an informative circulating free DNA test result confirming the presence of an activating anaplastic lymphoma kinase (ALK) rearrangement.  9. Histological or cytological evidence.  9. Documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC and there is an informative circulating free DNA test result confirming the presence of an activating anaplastic lymphoma kinase (ALK) rearrangement.  4. The patient has not previously received any ALK inhibitor for the advanced NSCLC indication unless 1st line treatment with Indiatinib, alectinib, certinib or crizotinib has had to be stopped within 6 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or  1. The patient has never previously received an ALK inhibitor or  1. The patient has never previously received and ALK inhibitor or  1. The patient has never previously received and ALK inhibitor or  1. The patient has never previously received and alk inhibitor or  1. The patient has previously received and such alk inhibitor or  1. The patient has previously received and such alk inhibitor or  1. The patient has previously received and such alk inhibitor or  1. The patient has previously received and such alk inhibitor or  1. The patient has previously received and such alk inhibitor or  1. The	No	TA670	27-Jan-21	27-Apr-21
CABA1	Cabazitaxel	Cabazitaxel for hormone-relapsed metastatic prostate cancer treated with docetaxel	1.1 Brigatnib will otherwise be used as set out in its Summary of Product Characteristics (SPC).  1.1 confirm that an application has been made and the first cycle of systemic anti-cancer therapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2.1 confirm the patient has hormone-relapsed metastatic prostate cancer.  3.1 confirm the patient has received 225mg/m/sq or more of docetaxel and the disease has progressed during or after docetaxel chemotherapy.  4.1 confirm cabazitaxel is to be prescribed in combination with prednisone or prednisolone.  5.1 confirm the patient has an Eastern Cooperative Oncology Group (ECOS) performance status of 0 or 1.  6.1 confirm the patient has been informed that treatment with cabazitaxel will be stopped if the disease progresses or after a maximum of 10 cycles (whichever happens first).  7.1 confirm the licensed dose and frequency of cabazitaxel will be used.	Yes	TA391	25-May-16	25-May-16

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with the combination of caboxantinib plus nivolumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has unresectable locally advanced or metastatic renal cell carcinoma (RCC) which has either a clear cell component or is one of the types of RCC as indicated below.  Please indicate below which RCC histology applies to this patient:  - RCC with a clear cell component or - Papillary RCC or - Chromophobe RCC or - Chromophobe RCC or - Chromophobe RCC or - Muclinous tubular and spindle cell RCC or - Muclinous tubular and spindle cell RCC or - XP11 translocation RCC or - XP11 translocation RCC or - Unclassified RCC				
			3. The patient has advanced RCC and the patient's disease is in the intermediate or poor risk category as assessed by the International Metastatic RCC Database Consortium (IMDC) system which scores 1 point for each of the 6 factors listed below – a score of 1 ori indicates good risk disease, a score of 1-2 indicates intermediate risk and a score of 3-6 denotes poor risk.  The IMDC factors are:  - less than 1 year from time of initial diagnosis of RCC to now - a Karmofsky performance status of <80% - the haemoglobin level is less than the lower limit of normal - the corrected calcium level is 2-25.mmol/L - the platelet count is greater than the upper limit of normal - the absolute neutrophil count is greater than the upper limit of normal.  Please indicate below whether the patient is in the intermediate or poor risk prognostic group: - intermediate risk disease (IMDC score of 3-6) Note: cabozantinib plus nivolumab is not approved for patients with good risk RCC.				
CABNIV1_v1.0	Cabozantinib in combination with nivolumab	For use in treatment-naïve patients with intermediate or poor risk advanced renal cell carcinoma for whom combination treatment with either involumab plus ipilimumab or lenvatinib plus pembrolizumab would otherwise be suitable where the following criteria have been met:		No	TA964	10-Apr-24	09-Jul-24
			6. The patient has a Karnofsky performance status of at least 70 (ie an ECOG performance score of 0 or 1).  7. The patient has no symptomatic brain metastases or leptomeningeal metastases currently requiring steroids for symptom control.  8. The patient is to be treated with cabozantinib until loss of clinical benefit or excessive toxicity or withdrawal of patient consent, whichever is the sooner.  Note: there is no stopping rule as to the maximum treatment duration of the cabozantinib part of this indication.  Note: if cabozantinib is permanently discontinued on account of toxicity, treatment with nivolumab can be continued as monotherapy as long as there is no evidence of progressive disease and the patient has not already completed 2 years of treatment with nivolumab.				
			9. The patient will receive the licensed dose, frequency, and route of nivolumab for this indication, as shown below  • subcutaneously – at a dose of 600mg every 2 weeks, or 1200mg every 4 weeks  • Intravenously – at a dose of 240mg every 2 weeks, or 480mg every 4 weeks  10. The patient is to be treated with nivolumab until loss of clinical benefit or excessive toxicity or withdrawal of patient consent or completion of a total treatment duration of 2 calendar years*, whichever occurs first.	-			
			*2 calendar years of treatment is defined as a duration of treatment which does not have any cycles of nivolumab in the period commencing on or after a date which is 2 years after the date of first nivolumab treatment.  Note: If nivolumab is permanently discontinued on account of toxicity, treatment with cabozantinib can be continued as monotherapy as long as there is no evidence of progressive disease.  11. When a treatment break of more than 12 weeks beyond the expected 2- or 4-weekly cycle length is needed, I will complete a treatment break approval form requesting a restart of treatment. This must be approved before nivolumab and/or cabozantinib is re-commenced				
			12. If the disease progresses on the cabozantinib plus nivolumab combination and further systemic therapy is appropriate, the next line of treatment will be chosen from those options which are routinely commissioned ie for the next line of systemic therapy, there will be use of one choice of the following (mainly incorporating TX options which have multiple modes of action): the currently commissioned 2nd line options of suntinib to lenvatinib price severolimus or everolimus monotherapy or the currently commissioned 1st line options of suntinib list lil on label as 2nd line treatment) or pazopamic (off label as 2nd line treatment) or pazopamic (off label as 2nd line treatment) or two actions of the currently commissioned 1st line options of suntinib list lil on label as 2nd line treatment) or pazopamic off label as 2nd line treatment).  13. Cabozantinib and nivolumab will be otherwise prescribed and administered as outlined in their respective Summary of Product Characteristics (SPCs).	-			

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is made by and the first cycle of systemic anti-cancer therapy with cabozantinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy				
			2. This patient has a confirmed histological diagnosis of medullary thyroid carcinoma				
			3. The patient has either metastatic disease or inoperable locally advanced disease				
			4. The disease is progressive and is either symptomatic or imminently likely to become symptomatic	1			
CABO1	Cabozantinib	The treatment of medullary thyroid cancer where all the following criteria are met:	5. The patient is treatment naïve to both cabozantinib and vandetanib unless the patient has had to discontinue vandetanib within 3 months of starting vandetanib because of toxicity (i.e. there is vandetanib toxicity which cannot be managed by dose delay or dose modification) and there has been no disease progression whilst on vandetanib.	Yes	TA516	28-Mar-18	26-Jun-18
		9	6. The patient has an ECOG performance status of 0 or 1 or 2.				
			7. Cabozantinib is to be continued as long as clinical benefit is observed or until there is unacceptable toxicity or patient choice to stop treatment				
			8. A formal medical review as to whether treatment with cabozantinib should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment	1			
			9. No treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve)				
			10. Cabozantinib is to be otherwise used as set out in its Summary of Product Characteristics				
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with cabozantinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy				
			2. The patient has a histologically- or cytologically-proven diagnosis of renal cell carcinoma (RCC) which either has a clear cell component or is one of the types of RCC as indicated below. Please indicate below which RCC histology applies to this patient:  - RCC with a clear cell component or - papillary RCC or - chromophobe RCC or - collecting duct RCC (Bellini collecting duct RCC) or - medullary RCC or - mucinous tubular and spindle cell RCC or - multilocular cystic RCC or - with respective response respons				
			3. The patient has either metastatic disease or inoperable locally advanced disease				
			4. The patient has previously received at least 1 vascular endothelial growth factor (VEGF)-targeted systemic therapy and has not been previously treated with cabozantinib.				
		The treatment of previously treated	Note: the patient may also have received prior treatment with an anti-PD-1, anti-PD-L1, anti-PD-L2, anti-CD137, or anti-Cytotoxic T-lymphocyte-associated antigen-4 (CTLA-4) antibody for renal cancer.				
CABO2	Cabozantinib	advanced renal cell carcinoma where the	5. The patient has progressed on previous treatment or within 6 months of most recent dose of VEGF inhibitor	Yes	TA463	08-Nov-17	08-Nov-17
		following criteria are met:	6. The patient has a performance status of 0 or 1				
			7. If the patient has brain metastases then these have been treated and are stable				
			8. Cabozantinib is to be continued until loss of clinical benefit or unacceptable toxicity or patient choice to stop treatment or cabozantinib can be stopped with a planned treatment break following the protocol used in the STAR trial.				
			Note: following 24 weeks of continuous cabozantinib therapy, and if there is no evidence of disease progression on therapy, patients and clinicians may choose to stop treatment for a planned drug free interval/treatment break and then restart cabozantinib on disease progression as per the STAR trial design.				
			Note: all patients who undergo planned treatment breaks must have regular clinical and radiological assessments and then have the option of restarting cabozantinib on disease progression.				
			Note: if the patient benefits from restarting after the first planned treatment break, they can take further planned treatments breaks following the same strategy, i.e. after a further 24 weeks on treatment.  Ref for the STAR trial: Brown JE, Royle KA, Gregory W, Ralph C, Maraveyas A, Din O et al. 'Temporary treatment cessation versus continuation of first-line tyrosine kinase inhibitor in patients with advanced clear cell renal cell carcinoms: (STAR): an open-label, non-inferiority, randomised, controlled, phase 2/3 trial.' The Lancet Oncology,2023, February 13 https://doi.org/10.1016/S1470-2045(22)00793-8.	3			
			9. A formal medical review as to whether treatment with cabozantinib should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment	1			
			10. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, a treatment break approval form will be completed to restart treatment unless the patient is following a planned intermittent treatment schedule as evidenced by the STAR trial and described above.	1			
			11. Cabozantinib will otherwise be used as set out in its Summary of Product Characteristics (SPC).	1			

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
CABO3	Cabozantinib	The treatment of treatment-naïve to vascular endothelial growth factor (VEGF)-targeted therapy and with intermediate or poor risk advanced renal cell carcinomawhere the following criteria are met:	1. This spatication is being made by and the first cycle of systemic anti-cancer therapy.  2. This patient has a histologically or cyclogically proven disposition from click carcinoma (ICC) which either has a dear cell component or is one of the types of RCC as indicated below.  Please indicate below which RCC histology agailes to this patient:  - RCC with a class cell component or - pupility RCC or - chromophable	Yes	TA542	03-Oct-18	01-Jan-19
CABO4	Cabozantinib	For the second line of tyrosine kinase inhibitor systemic therapy of Child-Pugh A locally advanced or metastatic hepatocellular carcinoma previously treated with sorafenib where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with cabozantinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has been previously treated with sorafenib for locally advanced or metastatic hepatocellular carcinoma.  3. The patient currently has Child-Pugh liver function class A.  4. The patient has an ECOS performance status of 0 or 1.  Note: NICE has not recommended cabozantinib in patients with an ECOS performance status of 2 or more.  5. The only other TKI with which the patient has been previously treated is sorafenib unless regorafenib has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of diseases progression.  6. The patient has not been previously treated with cabozantinib.  7. Cabozantinib is to be used only as monotherapy.  8. Cabozantinib is to be used only as monotherapy.  9. A formal medical review as to whether treatment with cabozantinib should continue or not will be scheduled to occur no later than by the end of the 2nd month of therapy.  10. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, a treatment break form will be completed to restart treatment, including an indication as appropriate if the patient had an extended break because of COVID 19.	Yes	TA849	14-Dec-22	14-Mar-23

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
CAP1	Capivasertib in combination with fulvestrant	Capivasertib in combination with fulvestrant for hormone receptor-positive, HER2-negate, locally advanced or metastatic breast cancer in patients previously treated with a CDKA/6 inhibitor and an aromatase inhibitor where the following criteria have been met:	1. This application for capital active therapy. 2. The patient has included the patient is a final active therapy. 3. The patient has included the patient is a final active therapy. 4. The patient has retained the patient is a final active therapy. 5. The patient has retained the patient is a final active therapy. 5. The patient has retained and accordance and active the patient is a final active therapy. 5. The patient has retained and previously freedom and the patient is a manufaction or suppression with Lifetime and previously treated with a patient is a manufaction or suppression with Lifetime and previously treated with an aromatose inhibitor. 5. If the patient has retained and previously freedom and the patient is male, consideration has been given to administration of Lifetime and previously treated with an aromatose inhibitor. 5. If the patient is female and previously freedom and the patient is a undergoon contrain ablation or suppression with Lifetime and previously treated with an aromatose inhibitor. 5. The patient has required and previously freedom and the patient is male, consideration has been given to administration of Lifetime and previously freedom and previous active and previously freedom with an aromatose inhibitor. 5. The patient has been previously freedom with an aromatose inhibitor. 6. The patient has been previously freedom with an aromatose inhibitor. 6. The patient has submitted a case to NICE for consideration of clinical and cost effectiveness only in patients previously freedom with a company active and an aromatose inhibitor. This population is narrower than that in the consideration has been previously freedom with a final patient is submitted that an aromatose inhibitor. This population is narrower th		TA1063		

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
CAR1	Carfilzomib	The treatment of previously treated multiple myeloma where all the following crtieria are met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with carfilzomib plus dexamethasone will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a diagnosis of multiple myeloma.  3. The patient has relapsed or progressing disease.  4. The patient has relapsed or progressing disease.  4. The patient has relapsed or progressing disease.  4. The patient has received 1 and only 1 prior line of treatment and that the numbering of a line of treatment is in accordance with the international Myeloma Workshop Consensus recommendations for the uniform reporting of clinical trials (http://dio.org/10.1182/blood-2010-10-299487). A line of therapy is defined as one or more cycles of a planned treatment program. This may consist of one or more planned cycles of single-agent therapy or combination therapy, as well as a sequence of treatments administered in a planned manner (eg induction chemotherapy/chemotherapies if followed by stem cell transplantation then maintenance is considered to be 1 line of therapy is of therapy is interrupted by a need for additional treatment for the disease.  Note: the use of carfilzomib in combination with dexamethasone in patients who have had 1 and only 1 prior line of therapy is because of NICE's specific recommendation for routine commissioning. The use of carfilzomib in combination with dexamethasone in patients who have had 1 and only 1 prior line of therapy is because of NICE's specific recommendation for routine commissioning. The use of carfilzomib in combination with dexamethasone in the 2-or more prior line patient groups is not permitted.  5. One of the following options applies as to any previous systemic therapy with bortezomib for this patient:  - the patient has not received any previous treatment with thortezomib or - the patient has not received any previous treatment with the bortezomib pare of state treatment and there has been at least a 6-month prote	Yes	TA657 (previously TA475)	18-Nov-20	17-Oct-17
CAR2	<b>Carfilzomib</b> in combination with lenalidomide and dexamethasone	For the treatment of previously treated multiple myeloma in patients who have had 1 prior line of systemic therapy where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with carfilzomib in combination with lenalidomide and dexamethasone will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a diagnosis of multiple myeloma.  3. The patient has relapsed or progressing disease.  4. The patient has received 1 and only 1 prior line of treatment and that the numbering of a line of treatment is in accordance with the international Myeloma Workshop Consensus recommendations for the uniform reporting of clinical trials (http://doi.org/10.1182/hlood-2010-10-29487). A line of therapy is defined as one or more cycles of a planned treatment program. This may consist of one or more planned cycles of single-agent therapy or combination therapy, as well as a sequence of treatments administered in a planned manner (eg induction chemotherapy)/chemotherapies if followed by stem cell transplantation and maintenance therapy is considered to be 1 line of therapy). A row line of therapy is modified to include other treatment agents (alone or in combination) as a result of disease progression, relapse or toxicity. A new line of therapy also starts when a planned period of observation off therapy is interrupted by a need for additional treatment for the disease.  Note: the use of carfilizomib in combination with lenalidomide and dexamethasone in patients who have had 1 and only 1 prior line of therapy is because of NICE's specific recommendation for this position in the myeloma treatment pathway. The use of carfilizomib in combination with lenalidomide and dexamethasone in the 2- or more prior line patients groups is not permitted.  5. The patient was treated with a bortezomib containing regimen as part of 1st line treatment and the patient responded to this bortezomib containing therapy.  Note: the company, when making its submission to NICE's subsequent on NICE's decination only in the group of patients who had responded t	No	TA695	28-Apr-21	27-Jul-21
			7. The patient has not been previously treated with carfilzomib. 8. Still me restament either included stem cell transplantation or not: 9. The patient has an ECOS performance status (PS) of 0 or 1 or 2. 10. The patient has an ECOS performance status (PS) of 0 or 1 or 2. 10. The patient will receive a maximum of 18 cycles of carfilzomib and that a patient continuing to respond after completing 18 cycles of carfilzomib plus lenalidomide plus dexamethasone will continue on treatment with lenalidomide plus dexamethasone without carfilzomib. 11. Carfilzomib will only be administered in combination with lenalidomide and dexamethasone and with no other systemic anticancer therapies. 12. Carfilzomib will only be administered in combination with lenalidomide and dexamethasone is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment or patient proceeds to stem cell transplant*, whichever is the sooner 12. Carfilzomib will henalidomide and dexamethasone is intended to be used for transplant ineligible patients after relapse or progression of first line therapy. Any patient receiving carfilzomib with lenalidomide and dexamethasone in this indication who subsequently becomes transplant eligible and is then able to proceed to transplant cannot resume treatment post-transplant as carfilzomib with lenalidomide and dexamethasone is not funded as maintenance therapy post-transplant.  13. A formal medical review as to whether treatment with carfilzomib plus lenalidomide plus dexamethasone should continue or not will be scheduled to occur no later than by the end of the 2nd month of therapy.  14. Where a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break form to restart treatment, including an indication as appropriate if the patient had an extended break.				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
CEM1	Cemiplimab	Cemiplimab monotherapy for the treatment of adult patients with locally advanced or metastatic cutaneous squamous cell carcinoma where the following treatment criteria have been met:	1. This application has been made by and the first cycle of systemic anti-cancer therapy with cemiplimab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, hepatitis and cutaneous reactions including Stevens-Johnson syndrome and took epidermal necrolysis.  3. The patient has a histologically or cyclologically-confirmed diagnosis of cutaneous squamous cell carcinoma.  4. The patient has either locally advanced disease or metastatic disease and is not a candidate for curative surgery or curative radiotherapy.  Pleaser ecorch here whether the disease as incally advanced disease with results in the patient not being a candidate for curative surgery or curative radiotherapy or metastatic disease with spread which results in the patient not being a candidate for curative surgery or curative radiotherapy or metastatic disease with spread that includes distant metastasis (eg lung, liver, bone etc)  5. The patient does not have a contra-indication to being treated with cemiplimab and that I am aware that immunocompromised patients were not included in the main cemiplimab clinical study: exclusion criteria in this study excluded any patient with a previous soil organ transplant or autoimnume disease which required systemic therapy with immunosuppressive agents within the previous 5 years or a history of pneumonitis within the last 5 years.  Cemiplimab should therefore be used with caution in immunosuppressed patients. By ticking 'yes' in the adjacent box you are stating that if cemiplimab is being administered to an immunocompromised patient, then you have evaluated and fully discussed with the patient the benefits and the risks of treatment with cemiplimab and the patient of the patient than the patient th	No	TA802	29-Jun-22	27-Sep-22

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slueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. I confirm that this application is made by and the first cycle of systemic anti-cancer therapy with ceritinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has histological or cytological evidence of NSCLC that carries an anaplastic lymphoma kinase (ALK) rearrangement based on a validated test <u>OR</u> there is documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC AND there is an informative circulating free DNA test result confirming the presence of an activating anaplastic lymphoma kinase (ALK) rearrangement. Please mark below on which basis the diagnosis of ALK positive NSCLC has been made in this patient:  - Histological or cytological evidence.  - Documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC and there is an informative circulating free DNA test result confirming the presence of an activating anaplastic lymphoma kinase (ALK) rearrangement				
CER1	Ceritinib	Certinin for anaplastic lymphoma kinase- positive advanced non-mall-cell lung cancer previously treated with crizotinib where the following criteria are met:	3. I confirm that the only TKI treatment that the patient has progressed on is 1st line circitorib.  Certitinib in this indication is only funded in patients who have been treated with and progressed on crizotinib as their sole TKI treatment.  4. I confirm that the patient has not been treated with 2nd line brigatinib after 1st line crizotinib unless the brigatinib had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression.  5. I confirm that the patient has not been previously treated with ceritinib.  6. I confirm that the patient has not been previously treated with ceritinib.  7. I confirm that ceritinib will be used only as monotherapy.  7. I confirm that the patient has an ECOG performance status of 0 or 1 or 2.  8. I confirm that the patient has not been previously treated with ceritinib.  9. I confirm that the patient will be treated with ceritinib until loss of clinical benefit or excessive toxicity or patient choice to discontinue treatment, whichever is the sooner.	No	TA395	22-Jun-16	20-Sep-16
			1.1. I confirm that ceritinib will be otherwise used as set out in its Summary of Product Characteristics  1. This application for ceritinib is being made by and the first cycle of systemic anti-cancer therapy with ceritinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has locally advanced or metastatic non-small cell lung cancer.  3. The patient has histological or cytological evidence of NSCLC that carries an anaplastic lymphoma kinase (ALK) rearrangement based on a validated test OR there is documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC AND there is an informative circulating free DNA test result confirming the presence of an activating anaplastic lymphoma kinase (ALK) rearrangement.  Please mark below on which basis the diagnosis of ALK positive NSCLC has been made in this patient:  - Histological or cytological evidence.  - Documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC and there is an informative circulating free DNA test result confirming the presence of an activating anaplastic lymphoma kinase (ALK) rearrangement				
CER2	Ceritinib	For anaplastic lymphoma kinase-positive advanced non-small cell lung cancer previously untreated with an ALK inhibitor where the following criteria have been met:	4. The patient has not previously received any ALK inhibitor unless 1st line alectinib or 1st line brigatinib or 1st line crizotinib has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression.  Please mark below which of the four scenarios applies to this patient:  - the patient has never previously received an ALK inhibitor or  - the patient has previously received alectinib as 1st line ALK-targeted therapy and this has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or  - the patient has previously received brigatinia a 1st line ALK-targeted therapy and this has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or  - the patient has previously received crizotinib as 1st line ALK-targeted therapy and this has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression.	No	TA500	24-Jan-18	24-Apr-18
			5. The patient is treatment-naive to 1st line cytotoxic chemotherapy-containing systemic treatment for this locally advanced or metastatic NSCLC indication.  Note: the only previous cytotoxic treatment allowed for patients to be treated with 1st line ceritinib is adjuvant or neoadjuvant chemotherapy given concurrently with radiotherapy.  6. The patient has an ECOG performance status of 0 or 1 or 2.  7. The patient either has no known brain metastases or if the patient has brain metastases, the patient is symptomatically stable prior to starting ceritinib.  8. Ceritinib will be used as monotherapy.  9. The patient will be treated until loss of clinical benefit or excessive toxicity or patient choice to discontinue treatment, whichever is the sooner  10. A formal medical review as to whether treatment with ceritinib should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment.  11. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment, including as appropriate if the patient had an extended break on account of Covid-19.				
			12. The prescribing clinician is aware that a) none of alectinib or brigatinib or crizotinib are to be used following disease progression on ceritinib as there is no current clear evidence to support treatment with any of these agents after disease progression on ceritinib and b) after disease progression on ceritinib, the only subsequent ALK inhibitor commissioned by NHS England is loriatinib.  13. Ceritinib will otherwise be used as set out in its Summary of Product Characteristics (SPC).				

lueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
CET4_v1.2	Cetuximab in combination with FOLFIRINOX/ FOLFIOXIRI (5- fluorouraci), irinotecan and oxaliplatin) chemotherapy	For chemotherapy-naive metastatic or locally advanced and inoperable colorectal cancer where the following criteria have been met:	This application is being made by and the first cycle of systemic anti-cancer therapy.  2. This patient has R&S wild syspe metastatic or locally advanced and inoproble colorectal cancer.  3. This patient has not necessed privates or toxicely devices controlled in the use of systemic anti-cancer therapy.  3. This patient has not necessed privates cyclotoc-chemotherapy for metastatic disease unless there has been use of previous necadijivant combination cytotocic chemotherapy for potentially resectable metastatic colorectal cancer. Please mark below whether the patient has had necadijivant chemotherapy or not:  1. the patient has not his perivous necadijivant cytotocic chemotherapy for potentially resectable metastatic colorectal cancer or  1. the patient has not his provision recombination in being used as either 1st line it restament for metastatic colorectal cancer or  2. Cetualizab in this FOLFRIRNOV/FOLFOXRII combination is being used as either 1st line it restament for metastatic colorectal cancer or an experiment of the patient has been treated with 1st line pembrolizumab for MS-H/dMMR disease.  Pease mark below in which line of therapy the patient is having cetualmia by 1st FOLFRIRNOV/FOLFOXRII combination is being used as either 1st line in treatment for metastatic colorectal cancer or as 2nd line treatment if treated with 1st line pembrolizumab for MS-H/dMMR disease.  Pease mark below in which line of therapy the patient is having cetualmab and the patient of the patient has not received prior for the patient with previous device and the patient of the patient has not received prior for the patient with previous device and patient of controlled cancer or as 2nd line treatment if treated with 1st line pembrolizumab or 1st line nivolumab which was previously wealable as an interior CVVI oxygina.  3. The patient has not received prior treatment with cetualmab panitumumab unless this was received as part of combination chemotherapy for potentially resectable metastatic disease.  3. The patient has not recei	Yes	TA439	Guidance 29-Mar-17	

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
CET1_v1.2	<b>Cetuximab</b> in combination with irinotecan-based chemotherapy	For chemotherapy-naive metastatic or locally advanced and inoperable colorecta cancer where all the following criteria are met:	1. This patient has RAS wild spee metastatic or locally advanced and inoperable colorectal cancer.  1. This patient has not received previous protocols treatment for metastatic diseases unless there has been use of previous neeadjuvant combination cytotoxic chemotherapy for potentially resectable metastatic colorectal cancer.  1. Please mark below whether the patient has had necadjuvant cytotoxic chemotherapy or not:  1. The patient has not hap perivous necadjuvant cytotoxic chemotherapy or not:  1. The patient has not hap previous necadjuvant cytotoxic chemotherapy or not:  1. The patient has not hap previous necadjuvant cytotoxic chemotherapy for potentially resectable metastatic colorectal cancer or  1. The patient has not hap previous necadjuvant cytotoxic chemotherapy for potentially resectable metastatic colorectal cancer or  2. The patient has not happened to the patient is having exclusionably but an international previous necessary of the patient patient is having exclusionably but an international previous necessary of the patient patie	Yes	TA439	29-Mar-17	27-Jun-17

ueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
CET2_v1.3	Cetuximab in combination with oxaliplatin-based chemotherapy	For chemotherapy-naive metastatic or locally advanced and inoperable colorectal cancer where all the following criteria are met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with returnab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient has RAS wild-type metastatic or locally advanced and inoperable colorectal cancer.  3. This patient has not received previous cytotoxic chemotherapy for metastatic colorectal cancer or the patient has not help previous cytotoxic chemotherapy for metastatic colorectal cancer or the patient has the had reconflowed the complete previous reconfigurant cytotoxic chemotherapy for metastatic colorectal cancer or the patient has been treated with previous reconfigurant cytotoxic chemotherapy for potentially resectable metastatic colorectal cancer or as 2nd line treatment if the patient has the high previous reconfigurant cytotoxic chemotherapy for potentially resectable metastatic colorectal cancer or as 2nd line treatment if the patient is having calculated by the previous reconfigurant power of the patient is having calculated by an onaligistatin-based combination chemotherapy.  4. Cetualmab in this oxaligistatin-based chemotherapy is being used as 2nd line treatment for metastatic colorectal cancer as 2nd line treatment if treated with 1st line pembrolizumab for MS-H/dMMR disease.  5. The patient has not received prior treatment with cetualmab prior patients and previously available as an interint COVID option  5. The patient has not received prior treatment with cetualmab proprior used an exact proprior treatment with cetualmab practical cancer or as part of combination neoadjuvant chemotherapy with the intention of resection if the metastates disease.  5. The patient has not received prior treatment with cetualmab prantinumab unless this was received as part of combination neoadjuvant chemotherapy with the intention of resection if the metastates disease in the patient is received an enadajovant cetualmab prantinumab containing combination neoadjuvant cetualmab prantinumab	Yes	TA439	29-Mar-17	27-Jun-17

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. An application has been made and the first cycle of systemic anti-cancer therapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a confirmed histological diagnosis of squamous cell carcinoma.  3. The patient has a primary tumour that originated in the oral cavity.  4. The patient has recurrent and/or metastatic disease.				
		Cetuximab in combination with chemotherapy for the first cytotoxic-	5. The patient has not received any previous cytotoxic chemotherapy for this recurrent/metastatic oral cavity tumour unless it was part of multimodality treatment for locally advanced disease and was completed more than 6 months previously.  6. The patient has not received any systemic therapy for this recurrent/metastatic oral cavity tumour or the only systemic therapy for this recurrent/metastatic oral cavity tumour has been with pembrolizumab monotherapy.  7. The treatment will be given with palliative intent.				
CET3_V1.1	Cetuximab	recurrent/metastatic squamous cell cancer of the head and neck only originating in	8. Cetuximab is to only be used in combination with a maximum of 6 cycles of platinum-based combination chemotherapy followed by single agent cetuximab as maintenance therapy.  9. The patient has received no previous treatment with cetuximab for head and neck cancer.  10. The patient has an ECOG performance status of 0 or 1.	Yes	TA473	31-Aug-17	31-Aug-17
		are met:	10. The patient has an ecolo periormanice status or our 1.  11. Cetuximab is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment, whichever is the sooner.  12. When a treatment break of more than 6 weeks beyond the expected 3- or 4-weekly cycle length is needed, a treatment break approval from will be completed to restart treatment.				
			13. Consideration has been to be given to administration of cetuximab 500mg/m² every 2 weeks (e.g. if chemotherapy is scheduled on a 4 week cycle or during the maintenance phase of single agent cetuximab therapy).				
		The treatment of relapsed/refractory	14. Cetuximab will be otherwise used as set out in its Summary of Product Characteristics.  1. Application made by and first cycle of systemic anti-cancer therapy to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy				
CLO1	Clofarabine	acute lymphoblastic leukaemia where all the following criteria are met:	2. Acute lymphoblastic leukaemia 3. Relapsed/ refractory disease with intent to use treatment to bridge to bone marrow transplant  1. This application for crizotinib is being made by and the first cycle of systemic anti-cancer therapy with crizotinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer	Yes	n/a - NHS England clinical policy	-	01-Apr-21
			therapy.  2. The patient has locally advanced or metastatic non-small cell lung cancer.  3. The patient has histological or cytological evidence of NSCLC that carries an anaplastic lymphoma kinase (ALK) rearrangement based on a validated test OR there is documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC AND there is an informative circulating free DNA test result confirming the presence of an activating anaplastic lymphoma kinase (ALK) rearrangement.  Please mark below on which basis the diagnosis of ALK positive NSCLC has been made in this patient:  - ibistological or cytological evidence.  - Documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC and there is an informative circulating free DNA test result confirming the presence of an activating anaplastic lymphoma kinase (ALK) rearrangement				
CRI1	Crizotinib	For anaplastic lymphoma kinase-positive advanced non-small cell lung cancer previously untreated with an ALK inhibitor where the following criteria have been	4. The patient has not previously received any ALK inhibitor for the advanced NSCLC indication unless 1st line alectinib or 1st line brigatinib or 1st line ceritinib has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or the patient was treated with adjuvant alectinib and had disease progression more than 6 months after completing treatment with adjuvant alectinib. Please mark below which of the four scenarios applies to this patient:  - the patient has never previously received an ALK inhibitor or  - the patient has previously received alectinib as 1st line ALK-targeted therapy and this has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or  - the patient has previously received brigatinib as 1st line ALK-targeted therapy and this has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or  - the patient has previously received ceritinib as 1st line ALK-targeted therapy and this has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or  - the patient has previously received ceritinib as 1st line ALK-targeted therapy and this has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression.  - the patient has previously received ceritinib as 1st line ALK-targeted therapy and this has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression.  - the patient has previously received teritinib as 1st line ALK-targeted therapy and this has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progr	No	TA406 TA422	28-Sep-16	28-Dec-16
		met:	5. Either the patient is naïve to 1st line cytotoxic chemotherapy-containing systemic treatment for this locally advanced or metastatic NSCLC indication or the patient received 1st line cytotoxic chemotherapy-containing treatment for locally advanced/metastatic non-small cell lung cancer at a time when the ALK status was not known and the patient has since received no further therapy.  Please mark which of these 2 scenarios below applies to this patient:  - the patient has not received any previous 1st line cytotoxic chemotherapy-containing systemic treatment for the locally advanced or metastatic NSCLC indication or  - the patient previously received 1st line cytotoxic chemotherapy-containing systemic treatment for locally advanced or metastatic NSCLC at a time when the ALK status was not known.  6. The patient has an ECOG performance status of 0 or 1 or 2.				
			7. The patient either has no known brain metastases or if the patient has brain metastases, the patient is symptomatically stable prior to starting crizotinib.  8. Crizotinib will be used as monotherapy.  9. The patient will be treated until loss of clinical benefit or excessive toxicity or patient choice to discontinue treatment, whichever is the sooner  10. A formal medical review as to whether treatment with crizotinib should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment.				
			11. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment.  12. The prescribing clinician is aware that a) alectinib is not to be used following disease progression on crizotinib as there is no current clear evidence to support treatment with alectinib after disease progression on crizotinib and b) after disease progression on crizotinib, the only subsequent ALK inhibitors commissioned by NHS England as next line therapy is a choice of brigatinib or ceritinib. c) after disease progression during treatment with adjuvant alectinib or within 6 months of completion of treatment with adjuvant alectinib, treatment with crizotinib is not commissioned 13. Crizotinib will otherwise be used as set out in its Summany of Product Characteristics (SPC).				

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
CRI3	Crizotinib	1st or subsequent line systemic therapy for ROS1-positive inoperable locally advanced/metastatic non squamous non- small cell lung cancer where the following criteria have been met:	1. I confirm that this application is made by and the first cycle of systemic anti-cancer therapy with crizotinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The patient has histological or cytological evidence of NSCLC that carries a ROS1 gene rearrangement based on a validated test OR there is documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC AND there is an informative circulating free DNA test result confirming the presence of a ROS1 gene rearrangement.  Please mark below on which basis the diagnosis of ROS1 positive NSCLC has been made in this patient:  -listological or cytological evidence.  - Documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC and there is an informative circulating free DNA test result confirming the presence of a ROS1 gene rearrangement  3. I confirm that this non squamous NSCLC carries a confirmed ROS1 gene rearrangement as demonstrated by an accurate and validated assay  4. I confirm that the patient has received no previous ROS1-targeted therapy  5. I confirm that the patient has received no previous ROS1-targeted therapy  5. I confirm that EITHER the patient has received no previous cytotoxic chemotherapy for locally advanced or metastatic disease  Note: NHS England has a strong preference for ROS1-positive patients to be treated with crizotinib as 1st line therapy for locally advanced/metastatic NSCLC though recognises that some patients have had to be treated with chemotherapy for urgent clinical reasons before the ROS1 result was known  6. I confirm that the patient has an ECOS performance status of 0 or 1 or 2  8. I confirm that the patient has no brain metastases or, if the patient has brain metastases, the patient is symptomatically stable prior to starting crizotinib  9. I confirm that the patient either has no brain metastases or, if the p	No	TA1021	04-Dec-24	03-Jan-25

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
DABTRA3	<b>Dabrafenib</b> in combination with trametinib	For the first line treatment of metastatic BRAF V600 mutation positive non-small cell lung cancer where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy.  2. The patient has a histologically confirmed diagnosis of non-small cell lung cancer (NSCLC).  3. The patient has histological or cytological evidence of NSCLC that contains a BRAF V600E mutation based on a validated test OR there is documented agreement by the lung MDT that the radiological appearances are in keeping with metastatic NSCLC AND there is an informative circulating free DNA test result confirming the presence of a BRAF V600E mutation.  Please mark below on which basis the diagnosis of BRAF V600E mutation positive NSCLC has been made in this patient:  - Histological or cytological evidence or  - Documented agreement by the lung MDT that the radiological appearances are in keeping with metastatic NSCLC and there is an informative circulating free DNA test result confirming the presence of a BRAF V600E mutation  4. The patient has metastatic non-small cell lung cancer.  5. I confirm that the patient is treatment naïve to BRAF and MEK inhibitors for the treatment of metastatic NSCLC.  6. I confirm that the patient has not received any previous systemic therapy for metastatic NSCLC.  Note: any prior adjuvant or neoadjuvant chemotherapy or immunotherapy for NSCLC does not count as previous systemic therapy in this regard.  7. The patient has an ECOG performance status of either 0 or 1 or 2.  Please enter below as to which ECOG performance status applies to this patient:  - ECOG PS 0 or  - ECOG PS 0 or  - ECOG PS 1 or  - ECOG PS 1 or  - ECOG PS 1 or  - ECOG PS 2  8. The patient bas no known brain metastases or if the patient has brain metastases, the patient is symptomatically stable prior to starting dabrafenib in combination with trametinib should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment.  10. A formal medical review as to how the combination of dabrafenib and trametinib is being tolerated and whether treatment with the combination of dabrafenib and trameti	Yes	TA898	14-Jun-23	12-Sep-23
DABTRA4	Dabrafenib (as Finlee*) in combination with trametinib (as Spexotras*)	For the treatment of paediatric patients aged 1-17 years with BRAF V600E mutation positive glioma where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with dabrafenib in combination with trametinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient is currently aged between 1 and 17 years.  3. The patient has a histologically confirmed diagnosis of either a low grade or a high grade glioma and that a BRAF V600E mutation has been confirmed to be present in whichever glioma type.  4. The patient either has a low grade glioma with a BRAF V600E mutation and requires systemic therapy or the patient has a high grade glioma with a BRAF V600E mutation and has received at least one prior radiation therapy and/or chemotherapy.  Please mark below which scenario applies to this patient:  - low grade glioma requiring first ever systemic therapy or  - lipid grade glioma having previously had ordinetrapy and chemotherapy or  - lipid grade glioma having previously had radiotherapy and have therapy or  - lipid grade glioma having previously had dinotherapy and chemotherapy only  5. The patient is either treatment naive to BRAF and MEK inhibitors for the glioma or the patient is currently receiving dabrafenib in combination with trametinib via a company compassionate access scheme and all treatment criteria on this form are fulfilled.  Please indicate below which option applies:  - No prior BRAF and MEK inhibitors for the treatment of glioma or  - the patient is currently receiving dabrafenib in combination with trametinib via a company compassionate access scheme and all treatment criteria on this form are fulfilled.  - The patient is currently receiving dabrafenib in combination with trametinib via a company compassionate access scheme and all treatment criteria on this form are fulfilled.  - The patient is currently receiving dabrafenib in combination with trametinib via a company compassionate access scheme and all treatment criteria on this form are fulfilled.  - The patient is currently receivi	No	TA977	29-May-24	27-Aug-24

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
DACO1	Dacomitinib	The treatment of untreated EGFR mutation-positive non-small-cell lung cancer where all the following criteria have been met:	1. This application is made by and the first cycle of systemic anti-cancer therapy with dacomitinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The patient has a histologically or cytologically confirmed diagnosis of non-small cell lung cancer (NSCLC) that is either stage III B or stage IV NSCLC  3. This patient's NSCLC has been shown to express an EGFR-activating mutation as demonstrated by an accurate and validated assay  4. The patient has received no previous EGFR-targeted therapy unless this has had had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression.  5. The patient has received no previous cytotoxic chemotherapy for locally advanced or metastatic non-small cell lung cancer  6. Dacomitinib will be used only as monotherapy  7. The patient has an ECOG performance status of 0 or 1  8. The prescribing clinician is aware of the potential drug interactions associated with dacomitinib therapy and the dose reductions or discontinuations required for the management of interstitial lung toxicity, diarrhoea and cutaneous toxicity.  9. The patient will be treated until loss of clinical benefit or excessive toxicity or patient choice to discontinue treatment, whichever is the sooner  10. Treatment breaks of up to 6 weeks are allowed but solely to allow toxicities to settle  11. Dacomitinib will be otherwise used as set out in its Summary of Product Characteristics (SPC)	No	TAS95	14-Aug-19	12-Nov-19

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is made by and the first cycle of systemic anti-cancer therapy with daratumumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a diagnosis of multiple myeloma.  3. The prescribing clinician understands that daratumumab in this indication is not funded for amyloidosis patients (with the exception of patients who have a proven diagnosis of myeloma and also have an associated diagnosis of amyloidosis) and that NHS funding for daratumumab is only for the specific multiple myeloma indication recommended by NICE.  Please tick box below:  - this patient has a proven diagnosis of primary amyloidosis  - this patient has a proven diagnosis of primary amyloidosis  - this patient has a proven diagnosis of primary amyloidosis  Note: For amyloidosis patients requiring systemic therapies, NHS England does fund treatments already in routine commissioning for myeloma. NHS England does not fund daratumumab in this indication for patients with amyloidosis unless they have a proven diagnosis of progressive myeloma and also an associated diagnosis of amyloidosis.  4. The patient has received 3 and no more than 3 prior lines of treatment and that the numbering of these lines of treatment is in accordance with the International Myeloma Workshop Consensus recommendations for the uniform reporting of clinical trials (http://doi.org/10.1182/blood-2010-10-299487). A line of therapy is defined as one or more cycles of a planned treatment program. This may consist of one or more planned cycles of single-agent therapy as the many as sequence of treatments administered in a planned manner (egi induction chemotherapy and stem cell treatpy) as modified to include other treatment agents (alone or in combination) as a result of disease progression, relapse or toxicity. A new line of therapy also starts when a planned course of therapy is modified to include other treatment agents (alone or in combination) as a result of disease				
DAR1	Daratumumab	The treating of relapsed and refractory multiple myeloma where all the following criteria are met:	5. The patient has responded to at least 1 of these 3 lines of treatment. 6. In relation to the immediately previous line of systemic therapy, the patient has: - documented relapse of disease after initial response or - refractory disease 7. The patient has been previously treated with a proteasome inhibitor. 8. The patient has been previously treated with an immunomodulatory agent. 9. Inave informed the CDF as to whether the patient has been treated with a previous stem cell transplant (SCT) or not: - Yes - previous SCT - No - previous SCT - 10. The patient is of performance status 0 or 1 or 2.	No	TA783	13-Apr-22	12-Jul-22
			- 0 - 1 - 2 - 2 - 11. The patient has not been previously treated with daratumumab or an anti-CD38 antibody unless they have been previously treated with daratumumab in which case the patient must have received the daratumumab as part of induction therapy pre-transplant and must have responded to that daratumumab-containing combination. The daratumumab-free period from previous therapy until now must be stated below.  Please enter below as to which scenario applies to this patient: - no previous treatment with daratumumab or - previous treatment with daratumumab in the transplant-eligible setting and disease responded to this. Please record the time since the start of the last cycle of daratumumab to now:				
			12. Daratumumab is only to be used as a single agent. It is not to be used in combination with other agents. The first administration of daratumumab can be given in split doses on different days if necessary.  13. A formal medical review as to whether treatment with daratumumab should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment.  14. Daratumumab is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment.  15. Where a treatment break of more than 6 weeks beyond the expected cycle length is needed, I will complete a treatment break approval form to restart treatment, including indicating as appropriate if the patient had an extended 16. Daratumumab will be otherwise used as set out in its Summary of Product Characteristics.				

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ilueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This aplication is being made by and the first cycle of systemic anti-cancer therapy with daratumumab in combination with bortezomib and dexamethasone will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a diagnosis of multiple myeloma.  3. The patient has a diagnosis of multiple myeloma.  3. The prescribing clinician understands that daratumumab in this indication is not funded for amyloidosis patients (with the exception of patients who have a proven diagnosis of myeloma with an associated diagnosis of amyloidosis) and that NHS funding for daratumumab is only for the specific multiple myeloma indication recommended by NICE.  Please tick box below:  - this patient has a proven diagnosis of primary amyloidosis.  - this patient has a proven diagnosis of primary amyloidosis.  - this patient has a proven diagnosis of progressive myeloma with an associated diagnosis of amyloidosis unless they have a proven diagnosis of progressive myeloma with an associated diagnosis of amyloidosis.  - The patient has received 1 and no more than 1 prior line of treatment and that the numbering of a line of treatment is in accordance with the International Myeloma Workshop Consensus recommendations for the uniform reporting of clinical trials (http://doi.org/10.1182/blood-2010-10-299487). A line of therapy is defined as one or more cycles of a planned treatment program. This may consist of one or more planned cycles of single-agent therapy or combination therapy, as well as a sequence of treatments administered in a planned manner (ie induction chemotherapy/chemotherapies when followed by stem cell transplantation is considered to be 1 line of therapy). A new line of therapy six when a planned course of therapy is modified to include other treatment agents, calone or in combination as a result of disease progression, relapse or toxidity, the exception to this being the need to attain a sufficient response for stem cell transplantation is				
DAR2	Daratumumab (in combination with bortezomib and dexamethasone)	For treating relapsed multiple myeloma in patients who have had only 1 line of therapy and are transplant ineligible where the following criteria have been met:	5. The patient responded to this 1-prior line of treatment (or if this patient received 2nd line ixazomib with lenalidomide and dexamethasone courtesy of Covid-related access IXA2CV, the patient must have responded to at least one of these 2 lines of therapy).  Note: the need for patients to have responded to their 1 prior line of treatment is as a consequence of the 1-prior subgroup chosen by Janssen for its submission to NICE for the appraisal of clinical and cost effectiveness of this daratumumab combination.  6. In relation to this 1-prior line of systemic therapy (or 2-prior in the case of patients accessing ixazomib with lenalidomide and dexamethasone via Covid-related access IXA2CV), the patient now has documented relapse of disease.  7. With respect to previous consideration of treatment with lenalidomide as part of previous therapy:  - this patient was treated with 1st line lenalidomide (either as 1st line therapy for transplant ineligible patients or as maintenance therapy in patients treated with stem cell transplantation as part of 1st line treatment) or  - the patient was treated with 2nd line ixazomib with lenalidomide and dexamethasone courtesy of the Covid-related access IXA2CV or  - treatment with 1st line lenalidomide in the transplant ineligible setting was considered unsuitable for this patient at the time or  - treatment with maintenance lenalidomide post stem cell transplantation was not available at the time or	Yes	TA897	06-Jun-23	04-Sep-23
			8. The patient has not been previously treated with daratumumab or an anti-CD38 antibody unless they have been previously treated with daratumumab in which case the patient must have received the daratumumab as part of induction therapy pre-transplant and must have responded to that daratumumab-containing combination. The daratumumab-free period from previous therapy until now must be stated below.  Please enter below as to which scenario applies to this patient:  - no previous treatment with daratumumab or - previous treatment with daratumumab in the transplant-eligible setting and disease responded to this. Please record the time since the start of the last cycle of daratumumab to now  9. With respect to current consideration of treatment with lenalidomide as part of 2nd line therapy:				
			- the patient has already been treated with lenalidomide with 1st line lenalidomide (either as 1st line therapy for transplant ineligible patients or as maintenance therapy in patients treated with stem cell transplantation as part of 1st line treatment) or received 2nd line lenalidomide as part of the Covid-related access IXA2CV to ixazomib with lenalidomide and dexamethasone - the patient is lenalidomide-naive but 2nd line treatment with lenalidomide is currently considered as unsuitable for this patient  10. The patient has either not been treated with high dose chemotherapy and stem cell transplantation or has been previously treated with high dose chemotherapy and stem cell transplantation or has been previously treated with high dose chemotherapy and stem cell transplantation or has been previously treatment with high dose chemotherapy and stem cell transplantation or previous treatment with high dose chemotherapy and stem cell transplantation or previous treatment with high dose chemotherapy and stem cell transplantation  11. the patient is of ECOS performance status 0 or 1 or 2.  Please tick one of the boxes below: - performance status 0 or - performance status 1 or				
			- performance status 2  12. Daratumumab is only to be used in combination with bortezomib and dexamethasone and that it is not to be used in combination with any other agents.  13. The dosage schedule of daratumumab will be for weekly treatment given in weeks 1.9 (a total of 9 doses), 3-weekly treatment in weeks 10 to 24 (a total of 5 doses) and 4-weekly treatment from week 25 onwards.  NHS England recommends that the subcutaneous formulation of daratumumab is used.  14. Daratumumab is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment.				
			15. A formal medical review as to whether treatment with daratumumab in combination with bortezomib and dexamethasone continues or not will be scheduled to occur at least by the end of the first 6 weeks of treatment.  16. When a treatment break of more than 6 weeks beyond the expected cycle length is needed, the prescribing clinician will complete a treatment break approval form to restart treatment.  17. Daratumumab will be otherwise used as set out in its Summary of Product Characteristics.				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
DAR3	Daratumumab in combination with bortezomib, thalidomide and dexamethasone	For induction and consolidation therapy of transplant-eligible multiple myeloma where the following criteria have been met:	1. This application is both being made by and the first cycle of systemic anti-cancer therapy.  It raise and and correlated in the use of systemic anti-cancer therapy.  It registers that should be a supported the state of systemic anti-cancer therapy.  It registers that should be a supported the state of systemic anti-cancer therapy for myeloma.  Note: this daratumumab indication is not funded for patients with primary amyloidosis.  Please confirm this by taking the box below.  It is patient does not have a diagnosis of primary amyloidosis.  It has patient does not have a diagnosis of primary amyloidosis.  It has patient as not previously received any systemic anti-cancer therapy for myeloma except for an emergency use of a short course of corticosteroids before this treatment.  It has patient is eligible for an autologous stem cell transplant after this induction therapy with the combination of daratumumab, bortezomib, thislidomide and desamethasone.  Sourtumumab will be given in combination with orteromib, talkidomide and desamethasone in the fow 28 day induction cycles pre-transplant and in the two 28 day cycles of post-transplant consolidation therapy.  Note: daratumumab is not funded for this transplant-eligible indication in combination with other anti-myeloma drugs.  It has patient is of ECOG performance status 0 or 1 or 2.  Please tick one of the boxes below:  - performance status 0 or  - performance status 1 or  - perfo	No	TA763	02-Feb-22	03-May-22

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
DAR4	<b>Daratumumab</b> in combination with lenalidomide and dexamethasone	For the treatment of newly diagnosed and treatment-naive patients with multiple myeloma who are INELIGIBLE for an autologous stem cell transplant where the following criteria have been met:	4. The patient is ineligible for an autologous stem cell transplant.  5. Daratumumab will only be given in combination with lenalidomide and dexamethasone and that it is not to be used in combination with any other agents.	No	TA917	25-Oct-23	23-Jan-24

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
DAR5	Daratumumab in combination with bortezomith cyclophosphamide and dexamethasone	For the treatment of newly diagnosed and treatment-naive patients with systemic immunoglobulin light chain anyloidosis (AL) where the following criteria have been met:	1. This application is both being made by and the first cycle of systemic anti-carect therapy with daratumumab in combination with bortezomis, cyclophosphamide and decamethasone will be prescribed by a consultant specialist specialist specialist and accreated in the use of systemic anti-carect therapy.  2. The patient has protocoly not received by any systemic anti-carect therapy for systemic light chain amyloidosis (AL).  3. The patient has protocoly not received by any systemic anti-carect therapy for systemic light chain amyloidosis (AL) of the patient is protocoly not received by any systemic anti-carect light chain amyloidosis (AL) of the patient is protocoly not received by a systemic anti-carect light chain amyloidosis (AL) from the patient system of the patient is systemic and the patient of the patient is systemic anti-carect light chain amyloidosis (AL) from the patient system is systemic anti-carect light chain amyloidosis (AL) and the patient is systemic anti-carect light chain amyloidosis (AL) from sof organ involvement or a system and the patient is systemic anti-carect light chain amyloidosis (AL). Forms of organ involvement could be cardiac, resul, hepatic, nervious system, gastrointestinal tract, lung and soft tissue. Passas is carect large involvement or 3 or more from organ involvement or 3 or more from organ involvement or 3 or more from organ involvement or 3. or more from	No	TA959	27-Mar-24	25-Jun-24

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
DAR5 (CONT)	Daratumumab in combination with bortezomib, cyclophosphamide and dexamethasone	For the treatment of newly diagnosed and treatment-naive patients with systemic immunoglobulin light chain amyloidosis (AL) where the following criteria have been met:	11. The the patient is of ECOG performance status 0 or 1 or 2. Please tick no or of the boxes below: - performance status 0 or - performance status 0 or - performance status 1 or - performance status 2  12. Daratumumab will only be given in combination with bortezomib, cyclophosphamide and dexamethasone and that it is not to be used in combination with any other agents.  13. The dosage schedule of daratumumab will be as follows: weekly treatment given in weeks 1-8 (a total of 8 doses in 2 x 4-weekly cycles) 2-weekly treatment in weeks 9-24 (a total of 8 doses in 4 x 4-weekly cycles) and from then on 4-weekly.  Note: the first administration of daratumumab can be given in split doses on different days if IV infusion is used instead of the preferred subcutaneous daratumumab formulation.  14. A maximum of 6 cycles of the combination of daratumumab plus bortezomib, cyclophosphamide and dexamethasone will be given unless there is development of progressive disease, unacceptable toxicity or patient choice to stop treatment.  15. Daratumumab monotherapy will continue to be given after completion of the combination therapy until whichever of the following events occurs first: the development of progressive disease, unacceptable toxicity or patient choice to stop treatment or after completion of a total 24 x 4-weekly cycles of daratumumab in combination with bortezomib, cyclophosphamide and dexamethasone.  Note: there is no funding for daratumumab after completion of a total 24 x 4-weekly cycles. It is therefore important that at the time of consenting, patients are informed of this maximum daratumumab treatment duration.	No	ТА959	27-Mar-24	25-Jun-24
			16. Hepatitis B virus screening has been recently done and that if positive hepatitis B viral serology is found, the patient will be monitored for hepatitis B virus reactivation as outlined in the daratumumab Summary of Product Characteristics.  17. A formal medical review as to whether treatment with daratumumab in combination with bortezomib, cyclophosphamide and dexamethasone continues or not will be scheduled to occur at least by the end of the second 4-weekly cycle of treatment.				
			18. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, the prescribing clinician will complete a treatment break approval form to restart treatment.  19. The National Amyloidosis Centre is auditing the outcomes of treatment-naïve patients commencing this daratumumab combination for light chain amyloidosis and details of this audit can be obtained by emailing Darren Foard (Clinical Nurse) Specialist) at a farenfaora@his.net  Note: NHS England strongly recommends participation in this audit which will provide real world evidence of this combination including data in patients with renal and cardiac involvement (some groups of which were excluded from the registration trial).  20. Daratumumab will be otherwise used as set out in its Summary of Product Characteristics.				

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
DARO1	Darolutamide in combination with androgen deprivation therapy (ADT)	For the treatment of non-metastatic hormone-resistant (castration-resistant) prostate cancer in patients who are at high risk of developing metastatic disease where the following criteria have been met	1. This application is being made by and the first cycle of systemic anti-cancer therapy with darolutamide will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient has a proven histological or cytological diagnosis of adenocarcinoma of the prostate without neuroendocrine differentiation or features of a small cell carcinoma.  3. This patient has non-metastatic prostate cancer as defined by recent imaging with conventional imaging with both a whole body isotope bone scan and a CT/MR scan of the chest, abdomen and pelvis.  Note: patients with the sole abnormality of pelvic lymph nodes measuring <2cm in short axis diameter and which are below the aortic bifurcation are eligible for darolutamide in this indication.  4. The patient has hormone-resistant (castrate-resistant) disease as defined by 3 rising PSA levels (after the nadir PSA level) and taken at least 1 week apart during androgen deprivation therapy.  5. The patient's serum testosterone level is <1.7 mmol/L on gonadotrophin releasing hormone aganist/antagonist therapy or after bilateral orchidectomy.  6. The current PSA level is ≥2ng/ml.  7. The patient is at high risk of developing metastatic disease as defined by a PSA doubling time of ≤10 months.  Please document the actual PSA doubling time in the box below:  8. The patient has an ECOS performance status of either 0 or 1 or 2.  9. The patient has an ECOS performance status of either 0 or 1 or 2.  9. The patient has an ECOS performance status of either 0 or 1 or 2.  9. The patient has an ECOS performance status of either 0 or 1 or 2.  9. The patient has an ECOS performance status of either 0 or 1 or 2.  9. The patient has not previously received any 2nd generation androgen receptor inhibitors (such as enzialutamide, apalutamide) or CYP17 enzyme inhibitors (such as abiraterone) unless the patient received apalutamide for non-metastatic hormone-resistant (history apalutamide patient received apalutamide for non-metastat	No	TA660	25-Nov-20	23-Feb-21

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
DARO2	Darolutamide in combination with docetaxel and androgen deprivation therapy (ADT)	For the treatment of patients with newly diagnosed metastatic hormone-sensitive prostate cancer where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy.  2. This patient either has a proven histological or cytological diagnosis of adenocarcinoma of the prostate or has presented with a clinical picture consistent with metastatic prostate cancer with both widespread bone metastases radiologically typical of prostate cancer and a serum PSA of 250 ng/ml.  3. This patient has newly diagnosed metastatic prostate cancer that is hormone sensitive and has currently received androgen deprivation therapy (ADT) for no longer than 12 weeks.  Please enter below as to which scenario applies to this patient:  - the patient has not yet received any ADT for metastatic prostate cancer or  - the patient has received no more than 12 weeks of ADT for metastatic prostate cancer  - the patient has received no more than 12 weeks of ADT for metastatic prostate cancer  - The patient is fit enough for docetaxel chemotherapy, has consented such treatment and has not yet commenced upfront docetaxel chemotherapy for metastatic hormone sensitive prostate cancer.  5. The patient has an ECOG performance status (PS) of O or 1.  Please enter below as to which ECOG performance status applies to this patient:  - ECOG OS 0  or  - ECOG PS 1  - To Darolutamide is being given in combination with both docetaxel and ADT.  8. The patient has not previously received any androgen receptor targeted agent such as enzalutamide or apalutamide or abiraterone unless the patient has progressive metastatic disease following completion of treatment with 2 years of ADT plus abiraterone with or without enzalutamide for high risk non-metastatic disease as part of the STAMPEDE trial (ISRCTN/78818544) and did not progress whilst on such treatment and the patient meets all the other criteria listed on this form.  Please mark below which of these 2 clinical scenarios applies to this patient: - the patient has not previously received any androgen receptor targeted agent - the patient has progressive metastatic disease aspart of the STA	No	TA903	21-Jun-23	19-Sep-23

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
DAS4	Dasatinib	imatinib-intolerant Philadelphia chromosome positive chronic phase chronic myeloid leukaemia in children where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with dasatinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has Philadelphia chromosome positive CML in chronic phase.  3. The patient has been previously treated with innatinib which had to be discontinued due to resistance or intolerance.  Please mark below whether the patient was resistant to or intolerant of imatinib: - resistant to innatinib or - intolerant of innatinib  4. The use of dasatinib has been discussed by the relevant multi-disciplinary team (IMDT) involved in chronic myeloid leukaemia (CML) decision making, which must include at least two consultants in the subspecialty with active and credible expertise in the relevant field of whom at least one must be a consultant paediatrician.  5. The patient is a child and I understand the Summary of Product Characteristics (SPC) states that "there is no experience with treatment of paediatric patients below 2 years of age" and "there is limited data in imatinib-resistant or intolerant paediatric patients below 6 years of age".  6. Treatment with dasatinib will be as monotherapy and with dosing appropriate to the tablet formulation or the oral suspension as described in the separate tablet and oral suspension Summaries of Product Characteristics (SPCs).  7. The prescribing clinician understands the SPC cautions that in paediatric patients after at least 2 years of treatment, treatment-related adverse events associated with bone growth and development were reported and close monitoring of growth in paediatric patients under dasatinib treatment is therefore recommended.  8. When a treatment break of more than 6 weeks beyond the expected cycle length is needed, I will complete a treatment break approval form to restart treatment, including indicating as appropriate if the patient had an extended break because of COVID19.  9. Dasatinib Will otherwise be used as outlined in the	No	As referenced in TA42S	21-Dec-16	21-Mar-17
DAS6	Dacatinih		1. I confirm that an application has been made by and the first cycle of systemic anti-cancer therapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. I confirm that the patient has chronic phase myeloid leukaemia  3. I confirm that the patient has received no prior treatment unless it was dasatinib received as part of the SPIRIT 2 trial for this indication and the patient meets all other criteria listed here*  *In March 2018 patients previously entered into the Spirit 2 trial and receiving free-of-charge supplies of dasatinib can transition to NHS commercial supply.  4. I confirm that invalinib is not appropriate for this patient and that this has been discussed and supported by the relevant MDT involved in CML decision making unless they are already receiving dasatinib as part of the SPIRIT 2 trial for this indication and the patient meets all other criteria listed here  5. I confirm that dasatinib will be used as outlined in the Summary of Product Characteristics (SPC).	No	TA426	21-Dec-16	21-Mar-17

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
DIN1	Dinutuximab beta		1. An application is being made by and the first cycle of systemic anti-cancer therapy with dinutuximab beta will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for hypersensitivity reactions, capillary leak syndrome, neuropathic pain, peripheral neuropathy and ocular toxicity.  3. The patient is currently aged 12 months or older and has histologically documented neuroblastoma according to the International Neuroblastoma Staging System (INSS).  4. The patient has high risk disease defined as either INSS stage 2, 3, 4 and 4s with MYCN amplification or INSS stage 4 without MYCN amplification and aged >12 months at diagnosis.  5. The patient achieved at least a partial response to induction chemotherapy (defined as whatever the sequence of therapies which subsequently led to myeloablative therapy and stem cell transplantation.  7. The patient memains free of disease progression following induction chemotherapy and stem cell transplantation.  8. The patient has not received prior treatment with an anti-GD2 antibody antibody unless they were treated with dinutusimab beta as part of induction therapy (as defined above) in the SIOPEN HR-NBL-2 or SIOPEN Pilot studies and all other treatment criterial listed on this form are fulfilled.  9. Dinutusimab beta is not being given in combination with interleuklin-2.  10. A formal medical review as to whether treatment with dinutusimab beta should continue or not and at what dose will be scheduled to occur at least by the end of the first cycle of treatment.  11. The patient will be treated until disease progression or excessive toxicity or completion of 5 cycles of therapy or patient/parent/guardian (as appropriate) choice to discontinue treatment, whichever is the sooner.  12. Treatment breaks of up to 6 weeks beyond the expected cycle length are allowed.  13. Dimutusmish beta will oth	No	TA538	22-Aug-18	20-Nov-18
DIN2	Dinutuximab beta	in patients aged 12 months and above and who have then both responded to intensive induction chemotherapy used to treat high risk 1st line patients and been treated with myeloablative therapy and stem cell transplantation where the following criteria are met:	1. An application has been made by and the first cycle of systemic anti-cancer therapy with dinutuximab beta will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for hypersensitivity reactions, capillary leak syndrome, neuropathic pain, peripheral neuropathy and ocular toxicity  3. The patient is currently aged 12 months or older and has histologically documented neuroblastoma according to the International Neuroblastoma Staging System (INSS)	No	TA538	22-Aug-18	20-Nov-18

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
DOS2	<b>Dostarlimab</b> in combination with platinum-containing chemotherapy (carboplatin and paclitaxel)	For the 1st line treatment of adult patients with mismatch repair deficient or microsatellite instability-high endometrial carcinoma who have recurrent or primary advanced disease and who are not candidates for potentially curative surgery or radiotherapy or chemoradiotherapy but are eligible for systemic therapy where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with dostarlimable in combination with carboplatin and packased will be prescribed by a consultant specialist specifically trained and accordingd in the use of systems can incancer therapy.  2. The prescribing clinical is fully warver of the management of, and the treatment modifications that may be required for, immune-related adverse reactions due to anti-PD-1 treatments including pneumonitis, colitis, nephritis, emocribing under the complexity of the properties of the prope	Yes	TA897	22-May-25	20-Aug-25

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ueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
DUR1_v1.2	Durvalumab	The treatment of PD-L1 21% positive locally advanced and unresectable nonsmall-cell lung cancer which has not progressed following concurrent platinum-based chemoradiotherapy where all the following criteria are met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with durvalumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, hepatitis and skin toxicity.  3. The patients as histologically or cyclologically-confirmed diagnosis of non-small cell lung cancer.  4. PD-L1 testing with an approved and validated test to determine the PD-L1 Tumour Proportion Score (TPS) has been done prior to this application and either the result demonstrates a PD-L1 score of 1% or more and the result is set out below or the besacretianed despite a clear intent and a reasonable attempt to do so.  Please document the actual PD-L1 TPS cannot be accurated that there is insufficient tissue for PD-L1 analysis and the Lung Cancer MDT has concluded and documented that the gaining of a further tumour sample is hazardous to the patient.  Note: durvalumab is not approved for use if the PD-L1 result is <1% or negative.  5. The patient has locally advanced and unresectable non small cell lung cancer which is either stage IIIA or stage IIIB or stage IIIC at the time of commencing concurrent chemoradiotherapy.  Please tick the correct box as to staging:  - stage IIIC disease or - stage IIIB disease or - stage	No	TA798	22-Jun-22	20-Sep-22
			12. The patient has not received prior treatment with an anti-PD-L3, anti-PD-L				
		a:	13. A formal medical review as to whether treatment with durvalumab should continue or not will be scheduled to occur at least by the end of the first 3 cycles of treatment.  14. Treatment breaks of up to 12 weeks beyond the expected cycle length are allowed but solely to allow any immune toxicities to settle.				
			14. Treatment dreads or up to 12 weeks beyond the expected cycle length are allowed but solerly to allow any immune toxicities to settle.  15. Durvalumab will be otherwise used as set out in its Summary of Product Characteristics (SPC).				

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Blueteq Form re	: Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
DUR2_v1.0	<b>Durvalumab</b> in combination with gemcitabine and cisplatin	For the 1st line treatment of patients with locally advanced or unresectable or recurrent or metastatic biliary tract cancer where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, hepatitis and skin toxicity.  3. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, hepatitis and skin toxicity.  3. The patients has histologically or cotylogically-confirmed diagnosis of adenocarcinoma of the biliary tract which comprises intrahepatic cholangiocarcinoma or extrahepatic cholangiocarcinoma or gall bladder carcinoma.  Please mark below which of these 3 sites of disease applies to this patient:  - intrahepatic cholangiocarcinoma  - earthepatic carcinoma  - gall bladder carcinoma  - gal	No	TA944	10-Jan-24	09-Apr-24

1. This application is being made by and the first cycle of systemic anti-cancer therapy with neoadjuvant durvalumab in combination with chemotherapy will be prescribed by a consultant specialist specifically trained and accredited in		started
The same of systems can be come which introduces an experience of the present the same which shortedgy applies to the papers.    Page 2016   Page 1   Page 2016   Page 2016	15-Jan-25	15-Apr-25

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
DUR4	<b>Durvalumab</b> In combination with etoposide plus either carboplatin or cisplatin	For the first-line treatment of adult patients with extensive-stage small cell lung cancer where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with durvalumab in combination with etoposide plus carboplatin or cisplatin will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinogathis, hepatitis and skin toxicity.  3. The patient has a histologically or cytologically determined diagnosis of small cell lung cancer (SCLC).  4. The patient has been staged as having extensive stage small cell lung cancer (SCLC).  5. The patient has not received previous systemic therapy for his/her extensive stage SCLC. Previous treatment with concurrent chemoradiotherapy for limited stage SCLC is allowed as long as therapy was completed at least 6 months prior to the diagnosis of recurrent and extensive stage disease.  6. The patient has an ECCG performance status score of 0 or 1.  7. The patient will be treated with a maximum of four 3-weekly cycles of durvalumab in combination with etoposide (80-100mg/m² IV on days 1-3 or its oral equivalent on days 2-3) plus either carboplatin (AUC 5 or 6 mg/ml/min) or cisplatin (7-8 80mg/m²).  8. On completion of durvalumab in combination with chemotherapy and in the absence of disease progression, treatment with durvalumab amaintenance monotherapy will continue until disease progression or symptomatic deterioration or unacceptable toxicity or withdrawal of patient consent, whichever occurs first.  9. The dosing of durvalumab will be at an intravenous dose of 1500mg given every 3 weeks in combination with chemotherapy and then maintenance intravenous 4-weekly durvalumab progression or symptomatical deterioration or unacceptable toxicity or withdrawal of patient consent, whichever occurs first.  11. The patient has no symptomatically active brain met	No	TA1041	19-Feb-25	20-Mar-25

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
DURG	<b>Durvalumab</b> in combination with tremelimumab	For first-line systemic treatment of adult patients with locally advanced or metastatic and/or unresectable hepatocellular carcinoma where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with durvalumab in combination with tremelimumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a diagnosis of hepatocellular carcinoma and that one of the following applies to the patient (please tick appropriate box below as to which option applies):  - either option 1 applies in which case a biopsy is deemed to be very high risk or technically not fessible in the patient and both the criteria a and b below are also both met:  - at the decision not to biopsy has been made and documented by a specialist MCC multi-disciplinary team meeting  and be the tumour meets the non-invasive diagnostic criteria of MCC as set out below*.  It is expected that option 2 will only apply in exceptional circumstances.  Please mark below which of these 2 clinical scenarios applies to this patient:  Option 1: the patient has a confirmed histological diagnosis of hepatocellular carcinoma or  Option 2: the patient has a confirmed histological diagnosis of hepatocellular carcinoma or  Option 2: the patient has a confirmed histological diagnosis of hepatocellular carcinoma or  Option 2: the patient has a confirmed histological diagnosis of hepatocellular carcinoma or  Option 2: the patient has a confirmed histological diagnosis of hepatocellular carcinoma or  Option 2: the patient has a confirmed histological diagnosis of hepatocellular carcinoma or  Option 2: the patient has a confirmed histological diagnosis of hepatocellular carcinoma or  Option 2: the patient has a confirmed histological diagnosis of hepatocellular carcinoma or  Option 2: the patient has optionable with the patient has not necessary to the patient has not necessary to the patient has not received previous systemic manual or the patient has not received previous systemic therapy for his/her hepatocellular carcinoma.  S. The patient has not received previous systemic therapy for hi	No	TA1090	19-Aug-25	17-Nov-25

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ELAC1	<b>Elacestrant</b> monotherapy	For the treatment of oestrogen receptor- positive, HER2-negative, locally advanced or metastatic breast cancer in patients previously treated with at least 12 calendar months of therapy with a CDK4/6 inhibitor-based combination where the following criteria have been met:	1. This application for elacestrant is being made by and the first cycle of elacestrant will be prescribed by a consultant specialitis specialitis application for elacestrant is being made by and the first cycle of elacestrant will be prescribed by a consultant specialitis specialitis application for elacestrant is a being region and the second and	No	TA1036	05-Feb-25	06-мау-25

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ENC1_v1.1	Encorafenib (in combination with binimetinib)	The treatment of unresectable stage III or stage IV BRAF V600 mutation positive malignant melanoma where the following criteria are met:	1. This application is made by and the first cycle of systemic anti-cancer therapy 2. This patient has a confirmed histological diagnosis of malignant melanoma. 3. This patient's cancer has been shown to contain a BRAF V600 mutation. 4. The patient has unresectable stage III or stage IV disease that has been staged according to the AJCC 8th edition 5. The patient is treatment naive to BRAF V600 and MEK inhibitors for malignant melanoma unless either the patient has previously received adjuvant dabrafenib and trametinib and did not progress during such therapy or has received a sufficient trial of dabrafenib plus trametinib for advanced disease and this has had to be stopped solely as a consequence of persistent dose-limiting toxicity and in the documented absence of disease progression.  Note: sequential treatment is not commissioned with dabrafenib plus trametinib and then on disease progression with encorafenib plus binimetinib.  6. The patient has sufficient ECOG performance status to tolerate treatment with the combination of encorafenib plus binimetinib.  7. Treatment with encorafenib in combination with binimetinib will be continued until loss of clinical benefit or unacceptable toxicity or withdrawal of patient consent unless the patient is enrolled in the DyNAMic clinical trial (trial reference CTA 21266/0255/001-0001) in which case an intermittent adaptive dosing schedule as guided by circulating tumour DNA levels can be used as per the trial protocol.  8. A formal medical review as to whether treatment with encorafenib in combination with binimetinib should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment  9. Where a treatment break of more than 6 weeks beyond the expected cycle length is needed, the prescribing clinician will complete a treatment break approval form to restart treatment, including indicating as appropriate if the patient had an extended break because of COVID 19.  Note: patients in the DyNAMic clinical trial (trial reference CTA 21266/0	No	TA562	27-Feb-19	28-May-19
ENC2_v1.2	Encorafenib in combination with cetuximab	For previously treated BRAF V600E mutation positive metastatic or locally advanced and inoperable colorectal cancer where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with encorafenib in combination with cetuximab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a histologically proven diagnosis of colorectal adenocarcinoma.  3. This patient's colorectal cancer has been shown to be of RAS wild type.  4. This patient's colorectal cancer has been shown to contain a BRAF V600E mutation.  5. The patient has failed one or two prior regimens for either metastatic or locally advanced and inoperable disease. Note: if the patient progressed through adjuvant chemotherapy or within 6 months of completing adjuvant chemotherapy, the patient can be classed as having received one line of treatment for metastatic disease.  Please note below whether the patient has been previously treated with one or two prior regimens for advanced/metastatic disease:  - One prior regimen  5. The has not received prior treatment with any BRAF inhibitor or MEK inhibitor unless this patient was treated with neoadjuvant encorafenib plus cetuximab prior to surgery for locally advanced but operable colon cancer within the FO/TROT 4 clinical trial (ISRCTN83842641).  Please mark below which of these 2 clinical scenarios applies to this patient:  - No prior treatment with any BRAF or MEK inhibitor  - Treated with neoadjuvant encorafenib plus cetuximab prior to surgery for locally advanced but operable colon cancer within the FO/TROT 4 clinical trial  - Treated with neoadjuvant encorafenib plus cetuximab prior to surgery for locally advanced but operable colon cancer within the FO/TROT 4 clinical trial  - Treated with neoadjuvant encorafenib plus cetuximab prior to surgery for locally advanced but operable colon cancer within the FO/TROT 4 clinical trial  - Treated with neoadjuvant encorafenib plus cetuximab prior to surgery for locally advanced but operable colon cancer within the FO/TROT 4 clinical trial  - Treated with neoadjuvant	No	TA668	06-Jan-21	06-Apr-21

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ENT2	Entrectinib	Entrectinib for ROS1-positive recurrent or locally advanced or metastatic non-small-cell lung cancer previously untreated with a ROS1 inhibitor therapy where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with entrectinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has histological or cytological evidence of NSCLC that carries a ROS1 gene rearrangement based on a validated test OR there is documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC AND there is an informative circulating free DNA test result confirming the presence of a ROS1 gene rearrangement.  Please mark below on which basis the diagnosis of ROS1 positive NSCLC has been made in this patient:  - Histological or cytological evidence.  - Documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC and there is an informative circulating free DNA test result confirming the presence of a ROS1 gene rearrangement  3. The patient has not previously received a ROS1 inhibitor.  Note: previous treatment with crizotinib is not allowed. The NICE recommendation and the entrectinib Summary of Product Characteristics both state that entrectinib is indicated in the treatment of patients who have not been previously treated with ROS1 inhibitors.  Please tick appropriately below as to whether the patient has been previously treated with systemic therapy for recurrent or locally advanced or metastatic NSCLC or  - the only systemic therapy was for recurrent or locally advanced or metastatic NSCLC and was with cytotoxic chemotherapy.  4. The patient has not been previously treated with entrectinib unless entrectinib has been received as part of any compassionate use scheme and the patient meets all the other criteria set out here.  5. Entrectinib will be used only as monotherapy.  6. The patient has no brain metastases or, if the patient has brain metastases, the patient is symptomatically stable prior to starting entrectinib.  8. The patient will	No	TA643	12-Aug-20	10-Nov-20
ENZ3	Enzalutamide in combination with androgen deprivation therapy (ADT)	For the treatment of patients with newly diagnosed metastatic hormone-sensitive prostate cancer where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with enzalutamide will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient either has a proven histological or cytological diagnosis of adenocarcinoma of the prostate or has presented with a clinical picture consistent with metastatic prostate cancer with both widespread bone metastases radiologically typical of prostate cancer and a serum PSA of ≥50 ng/mL.  3. This patient has newly diagnosed metastatic prostate cancer that is hormone sensitive and has either not been treated with docetaxel and has currently received androgen deprivation therapy (ADT) for no longer than 3 months or has been treated with docetaxel and has currently received and rogen deprivation therapy (ADT) for no longer than 3 months or has been treated with docetaxel and has currently received and not so the patients and the patient has not been treated with docetaxel and has currently received no more than 3 months of ADT (before starting an androgen receptor targeted agent) or - the patient has not been treated with docetaxel and has currently received no more than 3 months of ADT before starting an androgen receptor targeted agent) or - the patient has an ECOG performance status (PS) of 0 or 1 or 2.  5. The prescribing dinician has assessed this patient's status as regards receiving upfront docetaxel and have concluded that the patient completed planned docetaxel therapy or discontinued docetaxel before completion of planned treatment duration or should not have been treated with docetaxel or chose not to be treated with docetaxel. Plan the docetaxel Plan the docetaxel Plan to docetaxel and such as the patient to a planned treatment duration of 6 cycles of docetaxel - the patient with docetaxel and conscious planned treatment duration of 6 cycles of docetaxel - the patient townsenced docetaxer and discontinued docetaxel prior to completion of 6 cycles on account of	No	TA712	07-Jul-21	05-Oct-21
			7. The patient has not previously received any androgen receptor targeted agent unless the patient has received darolutamide, apalutamide or abiraterone for newly diagnosed metastatic hormone-sensitive prostate cancer which had to be stopped because of dose-limiting toxicity in the clear absence of disease progression and the patient meets all the other criteria listed here or the patient has progressive disease following treatment with 2 years of ADT plus abiraterone with or without encalutamide for high risk non-metastatic disease as part of the TSTAMPEDE trial (ISRTAMPEDE-1 trial and has not progressed whilst on such treatment and the patient meets all the other criteria listed on this form.  Please mark below which of these 6 clinical scenarios applies to this patient:  - the patient has not previously received any androgen receptor targeted agent - the patient has not previously received any androgen receptor targeted agent - the patient commenced darolutamide which had to be stopped because of dose-limiting toxicity in the clear absence of disease progression and the patient meets all the other criteria listed here - the patient commenced aburaterial which had to be stopped because of dose-limiting toxicity in the clear absence of disease progression and the patient meets all the other criteria listed here - the patient commenced aburaterone which had to be stopped because of dose-limiting toxicity in the clear absence of disease progression and the patient meets all the other criteria listed here - the patient commenced aburaterone which had to be stopped because of dose-limiting toxicity in the clear absence of disease progression and the patient meets all the other criteria listed here - the patient toxic stream the patient meets all the other criteria listed here - the patient commenced aburaterone which had to be stopped because of dose-limiting toxicity in the clear absence of disease progression and the patient meets all the other criteria listed here - the patient has stream the patient meets all				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with enzalutamide will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			2. This patient either has a proven histological or cytological diagnosis of adenocarcinoma of the prostate or has presented with a clinical picture consistent with metastatic prostate cancer with both widespread bone metastases radiologically typical of prostate cancer and a serum PSA of ≥50 ng/mL.				
			3. This patient has hormone-relapsed (castrate-resistant) metastatic prostate cancer.				
			4. The patient has no or only mild symptoms after androgen deprivation therapy has failed.				
			S. Chemotherapy is not yet indicated.				
ENZ4	Enzalutamide resistant) metastatic prostate cancer before chemotherapy is indicated where before chemotherapy is indicated where the patient has not been previously received any treatment with enzalutamide or darolutamide or apalutamide or abiraterone or	- the patient has not been previously received any treatment with enzalutamide or darolutamide or apalutamide or abiraterone or - the patient has previously received abiraterone for this same pre-chemotherapy indication in hormone-relapsed (castrate-resistant) prostate cancer but it was stopped within 3 months of it starting due to dose-limiting toxicity and in	Yes	TA377	27-Jan-16	26-Apr-16	
			7. The patient has an ECOG performance status (PS) of 0 or 1 or 2.	-			
			8. Enzalutamide is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment.				
			9. A formal medical review as to how enzalutamide is being tolerated and whether treatment with enzalutamide should continue or not will be scheduled to occur at least by the start of the third 4-weekly cycle of treatment.				
			10. Where a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment, including indicating as appropriate if the patient had an extended break because of COVID 19.				
			11. Enzalutamide is to be otherwise used as set out in its Summary of Product Characteristics.				
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with enzalutamide will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			2. This patient either has a proven histological or cytological diagnosis of adenocarcinoma of the prostate or has presented with a clinical picture consistent with metastatic prostate cancer with both widespread bone metastases radiologically typical of prostate cancer and a serum PSA of ≥50 ng/mL				
			3. This patient has hormone-relapsed (castration-resistant) metastatic prostate cancer.				
			4. The patient has been treated with docetaxel-containing chemotherapy and has progressed during or following treatment.				
ENZ5	Enzalutamide for the treatment of patients with hormone-relapsed (castrate- 5. One of the following applies to this patient as regards any previous use of 2nd generation receptor inhibitors (such as enzalutamide, darolutamide or apalutamide) or CYP17 enzyme inhibitors (such as abiraten resistant) metastatic prostate cancer with Please enter below as to which scenario applies to this patient:  Enzalutamide disease progression during or following:  the patient has not previously received any treatment with enzalutamide or darolutamide or abiraterone or	- the patient has not previously received any treatment with enzalutamide or darolutamide or apalutamide or abiraterone or - the patient has previously received abiraterone for this same post-chemotherapy indication in hormone-relapsed (castrate-resistant) prostate cancer but it was stopped within 3 months of it starting due to dose-limiting toxicity and	No	TA316	23-Jul-14	21-Oct-14	
		6. The patient has an ECOG performance status (PS) of 0 or 1 or 2.	1				
			7. Enzalutamide is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment.	1			
			8. A formal medical review as to how enzalutamide is being tolerated and whether treatment with enzalutamide should continue or not will be scheduled to occur at least by the start of the third 4-weekly cycle of treatment.				
		9. Wh	9. Where a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment, including indicating as appropriate if the patient had an extended break because of COVID 19.				
			10. Enzalutamide is to be otherwise used as set out in its Summary of Product Characteristics.	] ]			

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
EPC1	Epcoritamab	For the treatment of previously treated adult patients with diffuse large 8-cell lymphoma who have received 2 or more lines of systemic therapy which have included polatuzumab vedotin was contraindicated where the following criteria have been met:	In the speciation is being made by what the first special days are all receive therapy with a procramation more therapy will be prescribed by a consultant specialist specially specially included and accordance of the service of the special specia	No	TA954	06-Mar-24	O4-Jun-24

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ERD1	Erdafitinib	Erdafitinib for unresectable locally advanced or metastatic urothelial carcinoma which has a susceptible fibroblast growth factor receptor 3 (FGFR3) genetic alteration in patients previously treated with at least one line of therapy containing a PD-1 or PD-L1 inhibitor administered in the unresectable locally advanced or metastatic treatment setting where the following criteria have been met:	1. This application for endifitinis be being made by and the first cycle of systemic anti-cancer therapy with endificing build be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient is an adult with a histologically or optologically confirmed diagnosis of urothelial carcinoma.  Please also indicate below whether the urothelial carcinoma is of upper tract origin or the urothelial carcinoma is of upper tract origin or the urothelial carcinoma is of lower tract origin or the urothelial carcinoma is of lower tract origin or the urothelial carcinoma is of lower tract origin or the urothelial carcinoma has been step for FGR3 genomic alteration is positive:  - The patient is an indicate below which genetic alteration is positive:  - one of these FGR3 gene fusions: FGR8-TACC3 or FGR3-BAIAP2L10 or both a FGRR anticoma and a FGR3 gene fusion (FGR8-TACC3 or FGR3-BAIAP2L10 or both a FGRR anticoma and a FGR8 SEAC or SASC or	No	TA1062	12-May-25	09-Aug-2

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ERIB1	Eribulin	Eribulin for treating locally advanced or metastatic breast cancer after 2 or more	1. I confirm that an application has been made by and the first cycle of systemic anti-cancer therapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. I confirm that the patient has advanced breast cancer	Yes	TA423	21-Dec-16	21-Dec-16
		chemotherapy regimens	3. I confirm that the patient has has at least 2 prior chemotherapy regimens for advanced disease 4. I confirm the licensed dose and frequency of eribulin will be used.				
EVE1	Everolimus	Everolimus with exemestane for treating advanced breast cancer after endocrine therapy	1. I confirm that this application is being made by and the first cycle of systemic anti-cancer therapy of everolimus with exemestane will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. I confirm that the patient has ER +ve, HER2 –ve metastatic breast cancer  3. I confirm that the patient has no symptomatic visceral disease  4. I confirm that the patient has no symptomatic visceral disease  5. I confirm that the patient has had previous treatment with a non-steroidal aromatase inhibitor  6. I confirm that the patient has had no previous treatment with exemestane for metastatic breast cancer  7. I confirm the patient has received no more than one line of cytotoxic chemotherapy for the treatment of advanced breast cancer.  8. I confirm the licensed dose and frequency of everolimus will be used.	Yes	TA421	21-Dec-16	21-Dec-16
EVE5	Everolimus	Everolimus for advanced renal cell carcinoma after previous treatment	1. I confirm that an application has been made by and the first cycle of systemic anti-cancer therapy is to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. I confirm that the patient has biopsy proven renal cell carcinoma  3. I confirm that the patient has progressed during or after treatment with vascular endothelial growth factor targeted therapy  4. I confirm that the use of everoniums will be a sper the Summary of Product Characteristics (SPC)	Yes	TA432	22-Feb-17	23-May-17
EVEG	Everolimus	The treatment of unresectable or metastatic neuroendocrine tumours of pancreatic origin with disease progression where all the following criteria are met:	1. An application has been made by and the first cycle of systemic anti-cancer therapy is to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The patient has histopathologically proven well differentiated neuroendocrine tumour of pancreatic origin  3. The patient has unresectable or metastatic disease  4. The patient has exhibited disease progression in past 12 months  5. The patient has a performance status of 0-1  6. The patient has had no previous treatment with a mTOR inhibitor.  7. No planned treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve).*  8. Everolimus will otherwise be used as set out in its Summary of Product Characteristics (SPC).	Yes	TA449	13-May-17	26-Sep-17
EVE7	Everolimus	The treatment of unresectable or metastatic neuroendocrine tumours of gastrointestinal or lung origin with disease progression where all the following criteria are met:	1. An application has been made by and the first cycle of systemic anti-cancer therapy is to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The patient has histopathologically proven well differentiated neuroendocrine tumour of gastrointestinal or lung origin  3. The patient has surresectable or metastatic disease  4. The patient has no history of and no active symptoms to suggest a functional tumour  5. The patient has a seriformance status of 0-1  6. The patient has a performance status of 0-1	Yes	TA449	13-May-17	26-Sep-17

26-Nov-2025

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
FED1	Fedratinib	For the treatment of patients with myelofibrosis previously treated with ruxolitinib where the following criteria have been met:	1. This patient is an adult with a diagnosis of primary myelofibrosis (also known as chronic idiopathic myelofibrosis) or post polycythaemia vera myelofibrosis or post essential thrombocythaemia myelofibrosis.  Please enter below as to which type of myelofibrosis applies to this patient: - primary myelofibrosis or - post polycythaemia vera myelofibrosis or - post polycythaemia vera myelofibrosis or - post polycythaemia wera myelofibrosis or - post polycythaemia wera myelofibrosis or - post polycythaemia myelofibrosis or - post polycythaemia wera myelofibrosis or - post essential thrombocythaemia myelofibrosis - The patient has patient polycosis or not polyco	Yes	TA1018	20-Nov-24	18-Feb-25

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
FRU1	Fruquintinib	Fruquintinib for patients with either metastatic or locally advanced and inoperable colorectal cancer who have received 2 or more prior anticancer treatment regimens including fluoropyrimidine-, oxaliplatin- and irinotecan-based chemotherapies with or without anti-VEGF agents and/or anti-EGFR-based agents AND for whom the combination of triflurdine plus tipiracil and bevacizumab is unsuitable where the following criteria have been met:	1. This application is both being made by and the first cycle of systemic anti-cancer therapy.  2. The patient has a histologically confirmed diagnosis of adenocarcinoma of the colon or rectum.  3. The patient has been previously treated of metastatic or locally advanced and inoperable disease.  4. The patient has been previously treated or metastatic or locally advanced and inoperable disease.  5. The patient has been previously treated or metastatic or locally advanced and inoperable disease.  6. The patient has been previously treated with anti-EGFR-containing chemotherapy or not.  Please tick which option applies to this patient:	Yes	TA1079	23-Jul-25	21-Oct-25

26-Nov-2025

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
FUT1	Futibatinib	For the treatment of patients for locally advanced or metastatic cholangiocarcinoma which has a fibroblas growth factor receptor 2 gene fusion/rearnagement in patients with disease progression during or after previous systemic therapy where the following criteria have been met:	1. This application for futibatinib is being made by and the first cycle of systemic anti-cancer therapy.  2. The patient has a histologically or cytologically confirmed diagnosis of cholangiocarcinoma.  Please also indicate below whether the cholangiocarcinoma is of intrahepatic or extrahepatic origin: - the cholangiocarcinoma is of intrahepata corigin - the cholangiocarcinoma is of extrahepatic origin - the cholangiocarcinoma is of extrahepatic origin - the cholangiocarcinoma is of extrahepatic origin - the cholangiocarcinoma has been tested for fibroblast growth factor receptor 2 (FGFR2) gene fusion or rearrangement with a validated test and the result is positive.  4. The patient has unresectable locally advanced or metastatic disease.  5. The patient has been previously treated with systemic therapy for cholangiocarcinoma and the disease has progressed during or after such therapy.  Please also indicate whether the patient has received 1 or >= 2 lines of systemic therapy for cholangiocarcinoma - the patient has been previously treated with 1 line of systemic therapy for cholangiocarcinoma - the patient has been previously treated with 1 line of systemic therapy for cholangiocarcinoma - the patient has been previously treated with >= 2 lines of systemic therapy for cholangiocarcinoma - the patient has not ECOG performance status of 0 or 1.  7. The patient has not ECOG performance status of 0 or 1.  8. The patient has not previously received any specifically FGFR2-targeted therapy unless either the patient has received fultibatinib via a company early access scheme and the patient meets all the criteria set out on this form or pemigratinib monotherapy has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of progressive disease.  9. Futbatient has not been previously treated with a FGFR2-targeted therapy - the patient has not been previously treated with 1 in Graph and the patient: - the patient has not been previously treated with a FGFR	No	TA1005	11-Sep-24	10-Dec-24
			1.4. A first formal medical review as to whether treatment with fullibatinib should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment.  15. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, i will complete a treatment break approval form to restart treatment.  16. Fullibatinib will be otherwise used as set out in its Summary of Product Characteristics (SPC).				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
GEM1		Gemtuzumab ozogamicin as part of chemotherapy for previously untreated CD33 positive acute myeloid leukaemia in patients AGED 15 YEARS AND OVER where the following criteria are met:	1. This application is made by and the first cycle of systemic anti-cancer therapy with gemtuzumab ozogamicin will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The prescribing clinician is fully aware of the potential for gemtuzumab ozogamicin inducing hepatotoxicity including veno-occlusive liver disease/sinusoidal obstruction syndrome  3. This patient has a confirmed diagnosis of CD33-positive acute myeloid leukaemia but does NOT have acute promyelocytic leukaemia  4. The patient has previously untreated acute myeloid leukaemia  5. The patient is aged 15 years and over  Note: there is a separate application form for those patients who are aged less than 15 years  6. This patient has had cytogenetics performed  7. The result of the cytogenetics test has shown that the patient has one of the following (please tick appropriate box): - favourable risk stratification according to the 2017 EUN risk stratification OR - intermediate risk stratification according to the 2017 EUN risk stratification OR - the result of the cytogenetics test was unsuccessful OR - the result of the cytogenetics test is awaited and there is a clinical need for urgent systemic therapy to be commenced. If this is the case, it is mandatory that gemtuzumab ozogamicin will be discontinued as soon as cytogenetic results indicate adverse cytogenetics. Such discontinuation of gemtuzumab ozogamicin way be before all of the 1st cycle of induction treatment has been administered. Ticking the 'Need for urgent treatment before cytogenetics results indicate adverse cytogenetics. Such discontinuation of gemtuzumab ozogamicin will be stopped as soon as adverse cytogenetics are known.  8. The patient is fit for intensive induction chemotherapy  9. Gemtuzumab ozogamicin is to be given in combination with midoataurin (with either DA or FLA5-fida chemotherapy) for patients with a FLT3 mutation according to the trial protocol or the patient has been entered into the VICTOR clinical	No	TAS4S	14-Nov-18	12-feb-19
GEM2	Gemtuzumab ozogamicin		12. The use of gemtuzumab ozogamicin is exempt from the NHS England Treatment Break policy  1. An application has been made by and the first cycle of systemic anti-cancer therapy with gemtuzumab ozogamicin will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the potential for gemtuzumab ozogamicin inducing hepatotoxicity including veno-occlusive liver disease/sinusoidal obstruction syndrome  3. The patient has a confirmed diagnosis of C033-positive acute myeloid leukaemia but does NOT have acute promyelocytic leukaemia  4. The patient has previously untreated acute myeloid leukaemia  5. The patient has previously untreated acute myeloid leukaemia  5. The patient has previously untreated acute myeloid leukaemia  6. The post pubescent and if not going into a clinical trial will receive gemtuzumab ozogamicin at the dosage described in the results of the gemtuzumab ozogamicin COG AAMI.0531trial in children and reported in J Clin Oncol 2014; 32: 3023-3023 coil 1.01200/JCO.2014.53.5628  7. The result of the result of the gemtuzumab ozogamicin in this indication in people aged 15 years and over.  6. This patient has had cytogenetics performed  7. The result of the cytogenetics test has shown that the patient has one of the following (please tick appropriate box): favourable risk stratification according to the 2017 EUR risk stratification OR intermediate risk stratification according to the 2017 EUR risk stratification OR intermediate risk stratification according to the 2017 EUR risk stratification OR intermediate risk stratification according to the 2017 EUR risk stratification OR intermediate risk stratification according to the 2017 EUR risk stratification OR intermediate risk stratification according to the 2017 EUR risk stratification OR intermediate risk stratification according to the 2017 EUR risk stratification OR intermediate risk stratification according to the 2017 EUR risk stratificati	No	TA545	14-Nov-18	12-Feb-19

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with gilteritinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a proven diagnosis of acute myeloid leukaemia.				
			3. The patient has a FMS-like tyrosine kinase 3 (FLT3) mutation (internal tandem duplication [ITD] or tyrosine kinase domain [TKD]) as determined by a validated test.				
			4. The patient has relapsed/refractory FLT3 positive acute myeloid leukaemia.				
		For treating relapsed/refractory FLT3 mutation positive acute myeloid	5. The patient has not received previous systemic therapy with other FLT3 inhibitors (with the exception of sorafenib or midostaurin or quizartinib used in first-line therapy or in clinical trials in 1st line therapy).				
			6. The patient has an ECOG performance status (PS) of 0, 1 or 2.				
			7. Use of gilteritinib will be as monotherapy.				
GILT1	Gilteritinib	leukaemia in adults where the following criteria have been met:	8. Gilteritinib will be continued until disease progression or unacceptable toxicity or the time at which the patient is considered to be cured or until the patient receives a haematopoietic stem cell transplant whichever occurs first.	No	TA642	12-Aug-20	10-Nov-20
			9. The prescribing clinician understands that patients whose disease responds to gilteritinib and who then go on to have a haematopoietic stem cell transplant cannot restart gilteritinib as maintenance therapy after the transplant. This is as a consequence of the optimised NICE recommendation.				
			Note: patients who receive a stem cell transplant for FLT3 AML and who have not previously received treatment with gilteritinib cannot commence maintenance gilteritinib. Such patients can only receive gilteritinib if they relapse post SCT.				
			10. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for differentiation syndrome consequent to gilteritinib administration.				
			11. When a treatment break of more than 6 weeks beyond the expected cycle length is needed, I will complete a treatment break approval form to restart treatment, including indicating as appropriate if the patient had an extended break because of COVID 19.				
			12. Gilteritinib will be otherwise used as set out in its Summary of Product Characteristics (SmPC).				

ueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baselin fundin starte
			1. I confirm that this application is being made by and the first cycle of systemic anti-cancer therapy with glofitamab monotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			annument unergy.  2. Locatifirm that the patient has a histologically confirmed diagnosis of diffuse large B cell lymphoma (DLBCL) or transformed follicular lymphoma to DLBCL				
			The definition of DLBCL includes the following:				
			The definition of Deck Includes are notworing.  DIBCL not otherwise specified (NOS) [including germinal centre B-cell (GCB) and activated B-cell (ABC) subtypes]				
			primary mediastinal large B cell lymphoma				
			T cell rich B cell lymphoma  Epstein-Barr virus (EBV) positive DLBCL				
			intravascular large B cell lymphoma				
			double hit and triple hit high grade B cell lymphoma				
			Note: Primary CNS lymphoma, Burkitt lymphoma and plasmablastic lymphoma are NOT included for treatment with glofitamab.				
			Please record in the box below whether the patient has DLBCL according to one of the above types of DLBCL or has transformed follicular lymphoma:				
			- the patient has DLBCL according to one of the types within the above definition OR - the patient has transformed follicular lymphoma (TFL) to DLBCL				
			3. I confirm that the patient has DLBCL or TFL which has either relapsed following or is refractory to 2 or more lines of standard routinely commissioned systemic therapies and that within these 2 lines of therapy there has been treatment with an anti-CD20 regimen and an anthracycline-containing regimen.				
			Note: patients with TFL must have received 2 or more lines of systemic therapy given specifically for the transformed follicular lymphoma.				
			4. I confirm below the number of lines of systemic therapy that the patient has received for the treatment of DLBCL.				
			Note: induction chemotherapy prior to and then followed by stem cell transplantation counts as 1 line of systemic therapy. Similarly, bridging chemotherapy prior to and then followed by CAR T therapy counts as 1 line of systemic				
			therapy.  Note: patients who have had only 1 line of systemic therapy are not eligible for treatment with glofitamab.				
			Please record the number of lines of previous systemic therapy below:				
			- Peace rector are instance or lines of previous systems are tap years 2 previous lines OR				
			- 3 previous lines OR				
			- 4 or more previous lines  5. Londrime blow whether the patient has been previously treated with stem cell transplantation:				
		For the treatment of previously treated	- No previous stem cell transplantation OR				
	Glofitamab	adult patients with diffuse large B-cell	- Yes, previous stem cell transplantation				
GLO1	monotherapy	lymphoma who have received 2 or more lines of systemic therapy where the	6. I confirm below whether the patient has been previously treated with CART therapy and if so at which place in the treatment pathway:  - No previous CART therapy OR	Yes	TA927	17-Oct-23	16-No
		following criteria have been met:	- Yes, previous CAR T therapy as 2nd line therapy OR				
			- Yes, previous CAR T therapy as 3rd or more line of therapy  7. I confirm that the patient has not been previously treated with glofitamab unless either glofitamab monotherapy needs to be continued following EAMS access/a Roche compassionate access scheme or the patient received and				
			responded to no more than 3 cycles of glofitamab monotherapy used specifically as bridging treatment prior to 3rd or more line of CART therapy.				
			Note: glofitamab cannot be used as bridging therapy for 2nd line CART therapy.				
			Please record in the box below which of the following applies to this patient:				
			- no previous treatment with glofitamab OR - continuation of previous treatment with glofitamab monotherapy via EAMS and all other criteria on this form are fulfilled OR				
			- continuation of previous treatment with glofitamab monotherapy via a Roche compassionate access scheme and all other criteria on this form are fulfilled OR				
			- previous treatment with no more than 3 cycles of glofitamab monotherapy specifically used as bridging therapy prior to 3rd or more line CART therapy and the patient responded to this glofitamab bridging therapy				
			8. I confirm that the patient has not received any treatment with a bispecific antibody targeting both CD20 and CD3 other than glofitamab as specified above in criterion 7.				
			Note: use of glofitamab after previous treatment with epcoritamab is NOT commissioned.  9. I confirm that the patient has an ECOG performance status score of 0 or 1.				
			2. Tournin that the patient has an ecolo performance status score or or of 1.  Old Confirm that I am aware that a single does of binutuzumab 1000mg monotherapy is to be given on cycle 1 day 1 to mitigate the risk of cytokine release syndrome.				
			1.1. Loonfirm that with the exception of the single dose of obinitutzumab in cycle. I golfitamab is to degine on the sum of the control of the single dose of obinitutzumab in cycle. I golfitamab is to administered as monotherapy and not in combination with any other systemic therapies for lymphoma.				
			12. Lunderstand that I am aware that the dosing schedule of glofitamab in cycle 1 is 2.5mg on day 8 and 10mg on day 15, increasing to 30mg per cycle from cycle 2 day 1 onwards.				
			13.1 confirm that treatment with glofitamab monotherapy will be stopped at whichever of the following events occurs first: disease progression or unacceptable toxicity or withdrawal of patient consent or after a maximum of twelve				
			3-weekly cycles of glofitamab.				
			Note: once glofitamab is stopped after 12 cycles of treatment, it cannot be re-started.				
			14. I confirm that I and the treating team are familiar with the grading of cytokine release syndrome, its monitoring and management and the indications for use of tocilizumab and both I and the treating team have all undergone training in these clinical issues.				
			15. I confirm that I and the treating team are aware that the patient must be admitted overnight for at least the cycle 1 day 8 administration of glofitamab and potentially for further glofitamab infusions if grade 2 or greater cytokine release syndrome occurs with the previous glofitamab infusion.				
			16. I confirm that 1 dose of tocilizumab is immediately available should tocilizumab be required for the treatment of cytokine release syndrome and access to an additional dose of tocilizumab within 8 hours of the previous				
			tocilizumab must be ensured.				
			17. I confirm that a formal medical review as to whether treatment with glofitamab should continue or not will be scheduled to occur at least by the end of the first 6 weeks of treatment.				
		1	18. I confirm that when a treatment break of more than 6 weeks beyond the expected 3-weekly cycle length is needed, a treatment break approval form will be completed to restart treatment.				1

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
IBR5	lbrutinib	For the treatment of relapsed/ refractory mantle cell lymphoma in patients who have either only received 1 prior line of systemic therapy or been treated with 22 prior lines if 2nd line therapy was initiated before NICE's recommendation in January	1. This application has been made by and the first cycle of systemic anti-cancer therapy is to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a confirmed histopathological diagnosis of mantle cell lymphoma  3. The patient has previously been treated with one and only one prior line of rituximab-containing chemotherapy.  Note: Patients treated with more than 1 line of prior therapy are not eligible for treatment with ibrutinib.  4. The presence of relapsed/refractory mantle cell lymphoma with documented progression of disease during or following rituximab-containing 1st line systemic therapy.  5. The patient has never received any prior therapy with a BTK inhibitor (ibrutinib or zanubrutinib or another BTK inhibitor) unless the patient has suffered unacceptable toxicity on therapy with zanubrutinib without any evidence of disease progression and is transferring to treatment with ibrutinib.  Please enter below which of these scenarios applies to this patient:  - the patient is treatment-naïve to a BTK inhibitor or  - the patient has been receiving therapy with zanubrutinib but has suffered unacceptable toxicity without any evidence of disease progression and is transferring to treatment with ibrutinib	Yes	TA502	31-Jan-18	01-May-18
			6. Ibrutinib is to be used as a single agent 7. Ibrutinib is to be continued until disease progression, unacceptable toxicity or the patient's choice to stop treatment. 8. The patient's performance status is 0 or 1 or 2 9. The patient is not on concurrent therapy with warfarin. 10. The rescribing clinician is aware that ibrutinib has clinically significant interactions with cytochrome P450 enzyme 3A4 (CYP3A4) inhibitors and inducers as described in ibrutinib's Summary of Product Characteristics. 11. When a treatment break of more than 6 weeks beyond the expected cycle length occurs, the prescribing clinician will complete a treatment break approval form to restart treatment. 12. Ibrutinib will be otherwise used as set out in its Summary of Product Characteristics				
IBR9_v1.1	ibrutinib monotherapy	lbrutinib monotherapy for the treatment of patients with chronic lymphatic leukaemia which has a 17 pd election or TP53 mutation where the following criteria have been met:	1. This application for ibrutinib is being made by and the first cycle of this systemic anti-cancer therapy.  2. The patient has been diagnosed with chronic lymphatic leukaemia (CLL) or small lymphocytic lymphoma (SLL).  3. The patient has been diagnosed with chronic lymphatic leukaemia (CLL) or small lymphocytic lymphoma (SLL).  3. The patient has been tested for 17p deletion and preferably for TP53 mutation as well and the results are positive for either 17p deletion or TP53 mutation or both.  Please indicate the result of these tests below:  - positive for 17p deletion and not tested for TP53 mutation or  - positive for 17p deletion and not tested for TP53 mutation or  - positive for 17p deletion and positive for TP53 mutation or  - negative for 17p deletion and and TP53 mutation or  - positive for 10th 17p deletion and and TP53 mutation.  4. The patient has symptomatic disease which requires systemic therapy.  5. The patient has not received any previous BTK inhibitor therapy for CLL/SLL unless 1st line acalabrutinib or 1st line zanubrutinib has had to be stopped as a consequence of dose-limiting toxicity and in the clear absence of disease progression  - the patient has not received any previous BTK inhibitor therapy for CLL/SLL or  - the patient has not received any previous BTK inhibitor therapy for CLL/SLL or  - the patient has not received any previous BTK inhibitor therapy for CLL/SLL or  - the patient has not received any previous BTK inhibitor therapy for CLL/SLL or  - the patient has not received any previous BTK inhibitor therapy for CLL/SLL or  - the patient has not received any previous BTK inhibitor therapy for CLL/SLL or  - the patient has not received any previous BTK inhibitor therapy for CLL/SLL or  - the patient has not received any previous BTK inhibitor therapy for CLL/SLL or  - the patient has not received any previous BTK inhibitor therapy for CLL/SLL or  - the patient has not received any previous BTK inhibitor therapy for CLL/SLL or	Yes	TA429	25-Jan-17	25-Apr-17
			6. The patient has an ECOG performance status of 0 or 1 or 2.  7. Use of ibrutinib in this indication will be as monotherapy.  8. The prescribing clinician is aware that ibrutinib should not be administered concomitantly with warfarin or other vitamin K antagonists and that ibrutinib has clinically significant interactions with CYP3A4 inhibitors and inducers (see ibrutinib's Summary of Product Characteristics).  9. Ibrutinib is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment, whichever is the sooner.  10. A formal medical review as to whether treatment with ibrutinib should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment.  11. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment, including as appropriate if the patient had an extended break on account of Covid-19.  12. Ibrutinib will be otherwise used as set out in its Summary of Product Characteristics (SPC).				

	ug NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	baseline funding started
IBR10_v1.2 <b>Ibruti</b> i	Ibrutinib monotherapy for the treatment of patients with previously treated chronic lymphatic leukaemia where the following criteria have been met:	1. This application for inbrutinib is being made by and the first cycle of this systemic anti-cancer therapy with ibrutinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has been previously diagnosed with chronic lymphatic leukaemia (CLL) or small lymphocytic lymphoma (SLL).  3. The patient has been previously diagnosed with chronic lymphatic leukaemia (CLL) or small lymphocytic lymphoma (SLL).  3. The patient has been previously diagnosed with chronic lymphatic leukaemia (CLL) or small lymphocytic lymphoma (SLL).  3. The patient has been previously diagnosed with chronic lymphatic leukaemia (CLL) or small lymphocytic lymphoma (SLL).  4. The patient has been previously trained and not tested for TP53 mutation  4. The patient has previously from the patient has previously the for TP53 mutation  4. The patient has previously treated with systemic therapy for CLL/SLL  5. The patient has been previously treated with systemic therapy for CLL/SLL  6. The patient has been previously treated with systemic therapy for CLL/SLL and the acalabrutinib or anabunutinib has had to be discontinuously obleve house and observed with systemic dose-imiting toxicity and in the clear absence of disease progression or the patient has previously been treated with the SLI time combination of ibrutinib plus venetociax and was still in response on completion of treatment but has since relapsed and this application will be the first use of a BTK inhibitor since the 1st line combination of ibrutinib plus venetociax and was still in response on completion of treatment but has since relapsed and this application will be the first use of a BTK inhibitor since the 1st line combination of ibrutinib plus venetociax and was still in response on completion of treatment but has since relapsed of dose-imiting toxicity and in the clear absence of disease progression or the patient previously commenced analbrutinib for relapsed/refractory CLU/SLL and acalabr	Yes	TA429	25-Jan-17	25-Apr-17

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
IBR11	<b>ibrutinib</b> in combination with venetoclax	For the 1st line treatment of previously untreated chronic lymphatic leukaemia where the following criteria have been met:	1. This application for ibrutinib in combination with venetociax is being made by and the first cycle of ibrutinib plus venetociax will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anticancer therapy.  2. The patient has been diagnosed with chronic lymphatic leukaemia or small lymphocytic lymphoma.  3. The patient has been tested for 17p deletion and negative for 1793 mutation. Please indicate the result of these tests below:  Negative for 17p deletion and negative for 1793 mutation  Negative for 17p deletion and negative for 1793 mutation  Negative for 17p deletion and positive for 17p	No	TA891	31-May-23	started
			15. Ibrutinib and venetoclax will be otherwise used as set out in their respective Summary of Product Characteristics (SPCs).				

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
INO1	Inotuzumab ozogamicin	The treatment of relapsed/refractory Philadelphia positive and Philadelphia negative B cell precursor acute lymphoblastic leukaemia in ADULT patients where all the following criteria are met:	1. An application is being made by and the first cycle of systemic anti-cancer therapy with inotuzumab ozogamicin for each part of the treatment pathway will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing officians is fully ware of the risk factors for inotuzumab ozogamicin inducing hepatotosicity including veno-occlusive liver disease/sinusoidal obstruction syndrome and that this risk rises as the number of cycles administrater increases.  3. The patient has relapsed or refractory CD22 positive B cell precursor acute lymphoblastic leukaemia (ALL).  Please tek the appropriate box as to which type of ALL the patient has:  **Philadeliphia chromosome positive ALL in which case treatment with at least one TKI must have also failed  4. The patient has been previously treated with intensive combination chemotherapy as initial treatment with or without subsequent salvage chemotherapy or blinatumomab.  5. The patient has and activ:  **Note: there is a separate Blueteq form to be used for inotuzumab ozogamicin in this indication in children.  6. Inotuzumab ozogamicin will only be requested by and administered in either bone marrow transplant centres or in major haematological centres that regularly treat patients with relapsed/refractory ALL and who have regular ALL multi-disciplinary team meetings and close links with bone marrow transplant centres or in major haematological centres that regularly treat patients with relapsed/refractory ALL and who have regular ALL multi-disciplinary team meetings and close links with bone marrow transplant centres or in major haematological centres that regularly treat patients with relapsed/refractory ALL and who have regular ALL multi-disciplinary team meetings and close links with bone marrow transplant centres or in major haematological centres that regularly treat patients with relapsed/refractory ALL and who have regular ALL multi-disciplinary team meetings and close links with meeting	No	TA541	19-Sep-18	18-Dec-18
INO2	Inotuzumab ozogamicin	The treatment of relapsed/refractory Philadelphia positive and negative B cell precursor acute hympholalstic leukaemia in CHILD patients where all the following criteria are met:	treatment break approval form.  13. Inotuzumab zoogamicin will otherwise be used as set out in its Summary of Product Characteristics (SPC).  1. An application has been made by and the first cycle of systemic anti-cancer therapy with inotuzumab zoogamicin will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the risk factors for inotuzumab zoogamicin inducing hepatotoxicity including veno-occlusive liver disease/sinusoidal obstruction syndrome and that this risk rises as the number of cycles administered increases.  3. The patient has relapsed or refractory CD22-positive B cell precursor acute lymphoblastic leukaemia (ALL).  Please teck appropriate box as to which type of ALL the patient has:  *Philadelphia chromosome positive ALL in which case treatment with at least one second or third generation TKI must have also failed  4. The patient has been previously treated with intensive combination chemotherapy as initial treatment with or without subsequent salvage chemotherapy or blinatumomab  5. The patient is a child* and:  • is post pubscent or  • is pre-pubscent and will receive inotuzumab zoogamicin at the dosage described in the results of the inotuzumab zoogamicin trial in children and reported in Pediatric Blood Cancer 2014; 61: 369-372 doi: 10.1002/pbc.24721  *note there is a separate Blueteq form to be used for inotuzumab zoogamicin in this indication in adults.  6. Inotuzumab zoogamicin will only be requested by and administered in principal treatment centres  7. The use of the inotuzumab zoogamicin has been discussed at a multi-disciplinary team (MDT) meeting which must include at least two consultants in the subspecialty with active and credible expertise in the relevant field of whom at least one must be a consultant paediatricin. The MDT should include a paediatric pharmacist and other professional groups appropriate to the disease area  8. The patient has a performance status of 0	No	TA541	19-Sep-18	18-Dec-18

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
IV01_v1.0	Ivosidenib monotherapy	For the treatment of patients with locally advanced or metastatic cholangiocarriomas which has an isocitrate dehydrogenase-1 (IDH1) R132 mutation in patients with disease progression during or after previous systemic therapy and where the following criteria have been met:	1. This application for lovisidenib is being made by and the first cycle of systemic anti-cancer therapy with lovisidenib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a histologically or cytologically confirmed diagnosis of cholangiocarcinoma.  2. The patient has a histologically or cytologically confirmed diagnosis of cholangiocarcinoma.  3. The cholangiocarcinoma is of extrahepatic origin  3. The cholangiocarcinoma is not extrahepatic origin  4. The patient has unresectable locally advanced or metastatic disease.  5. The patient has unresectable locally advanced or metastatic disease.  6. The patient has been previously treated with systemic therapy for cholangiocarcinoma and the disease has progressed during or after such therapy. Such systemic therapy could have been in the adjuvant or neoadjuvant or advanced disease sestings.  9. Please also indicate whether the patient has received 1 or 22 lines of systemic therapy.  1. The patient has been previously treated with 1 line of systemic therapy for cholangiocarcinoma or  1. The patient has been previously treated with 2 lines of systemic therapy for cholangiocarcinoma  2. The patient has been previously treated with 2 lines of systemic therapy for cholangiocarcinoma  3. The patient either has no known brain metastases or if the patient has brain	No	TA948	31-Jan-24	30-Apr-24

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
IV02_v1.0	Ivosidenib in combination with azacitidine	For newly diagnosed and untreated adult acute myeloid feukaemia with an isocitrate dehydrogenesse-1 (IDH1) R132 mutation in patients who are not eligible for standard induction chemotherapy where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy.  2. The patient has newly diagnosed acute myloid feutaments (AML).  3. The patient has newly diagnosed acute myloid feutaments (AML).  4. The patient has newly diagnosed acute myloid feutaments (AML).  4. The patient has newly diagnosed acute myloid feutaments (AML).  4. The patient has newly diagnosed acute myloid feutaments (AML).  4. The patient has newly diagnosed acute myloid feutaments (AML).  4. The patient has the most recent bone marrow blist count:  2. Office of AML.  5. The patient has the most recent bone marrow blist count:  2. Office of AML.  5. The patient has the most recent bone marrow blist count:  2. Office of AML.  5. The patient has the most recent bone marrow blist count:  2. Office of AML.  5. The patient has the most recent bone marrow blist count:  2. Office of AML.  5. The patient has the most recent bone marrow blist count:  2. Office of AML.  5. The patient has the most recent bone marrow blist count:  2. Office of AML.  5. The patient has the most recent bone marrow blist count:  2. Office of AML.  5. The patient has the most recent bone marrow blist count.  5. Office of AML.  5. The patient has the most recent bone marrow blist count.  5. Office of AML.  5. The patient is fire for treatment should not be a stated in which will be	Yes	TA979	05-Jun-24	03-Sep-24

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
IXA1_V1.1	ixazomib with lenalidomide and dexamethasone	The treament of relapsed or refractory multiple myeloma where all the following criteria are met:	1. The patient has nestablished diagnosis of multiple myeloma.  2. The patient has an established diagnosis of multiple myeloma.  3. The prescribing efficies numberstand; that this confinitation of value diagnosis of multiple myeloma and who also have an associated diagnosis of amyloidosis, and that Net's flanding for sazomia is only for the specific myeloma indication recommended by NiCE.  4. The patient has an established diagnosis of multiple myeloma and who also have an associated diagnosis of amyloidosis, and that Net's flanding for sazomia is only for the specific myeloma indication recommended by NiCE.  4. This patient does not have a diagnosis of primary amyloidosis patients grower adaption of progressive myeloma and who also have an associated diagnosis of amyloidosis and this isazomib combination is being prescribed for the myeloma. Note: for primary amyloidosis patients requiring systems therapies, NiCE does fund other treatments already in routine commissioning for myeloma. NiEE does not fund this bazomib combination in this indication for patients with amyloidosis.  4. The patient has received 2 or 3 prior lines of treatment (i.e. no lines lists than 2 and no lines more than 3) and that the numbering of these lines for treatment is in accordance with the international Myeloma Workshop Consensus recommendations for the uniform epoprating of clinical trisk (Interplace) as a sequence of treatments already in routine commissioning for treatment is an accordance with the international Myeloma Workshop Consensus recommendations for the uniform epoprating of clinical trisk (Interplace) as a sequence of treatments almost deal diagnosis of amyloidosis.  4. The patient has received 2 or 3 prior lines of treatment (i.e. no lines lists than 2 and no lines more than 3) and that the numbering of these interplaces of treatments and the patient of the search of the uniform epoprating of clinical trisk (Interplace) and a security of the lines does not a planned prior of controllation of the patient the secure of th	Yes	TA870	22-Feb-23	23-May-23

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
LEN1	Lenalidomide in combination with dexamethasone	The 1st line treatment in transplant ineligible patients with multiple myeloma in whom thaildomide is contraindicated or who cannot tolerate thaildomide where the following criteria have been met:			TAS87	26-Jun-19	started
			S. The patient is of ECOS performance status 0 or 1 or 2.  Please tick no or of the boxes below: - performance status 1 or - performance status 1 or - performance status 1 or - performance status 2 or - serformance status 2 or				
LEN2	Lenalidomide in combination with dexamethasone	The 2nd line treatment in transplant ineligible patients with multiple myeloma previously treated with a 1st line bortezomib containing regimen where the following criteria have been met:	1. This application is made by and the first cycle of systemic anti-cancer therapy with lenalidomide in combination with dexamethasone will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a confirmed diagnosis of multiple myeloma.  3. The patient has been treated with a 1st line regimen which contained bortezomib.  5. The patient has been treated with a 1st line regimen which contained bortezomib.  5. The patient has received 1 and no more than 1 prior line of treatment and that the numbering of a line of treatment is in accordance with the International Myeloma Workshop Consensus recommendations for the uniform reporting of clinical trials (http://doi.org/10.1182/blood-2010-10-299487). A line of therapy is defined as one or more cycles of a planned treatment program. This may consist of one or more planned cycles of single-agent therapy or combination therapy, as well as a sequence of treatments administered in a planned manner (le induction chemotherapy/chemotherapies when followed by stem cell transplantation then maintenance is considered to be 1 line of therapy is modified to include other treatment agents (alone or in combination) as a result of disease progression, relapse or toxicity, the exception to this being the need to attain a sufficient response for stem cell transplantation to proceed. A new line of therapy also starts when a planned period of observation off therapy is interrupted by a need for additional treatment for the disease.	No	TAS86	26-Jun-19	24-Sep-19

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is made by and the first cycle of systemic anti-cancer therapy with lenalidomide in combination with dexamethasone will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			2. The patient has a confirmed diagnosis of multiple myeloma.				
			3. The patient is ineligible for stem cell transplantation				
			4. The patient has received at least 2 prior lines of treatment and that the numbering of a line of treatment is in accordance with the International Myeloma Workshop Consensus recommendations for the uniform reporting of clinical trials (http://doi.org/10.1182/blood-2010-10-299487). A line of therapy is defined as one or more cycles of a planned treatment program. This may consist of one or more planned cycles of single-agent therapy or combination therapy as well as a sequence of treatments administered in a planned manner (le induction chemotherapy/chemotherapies when followed by stem cell transplantation then maintenance is considered to be 1 line of therapy). A new line of therapy starts when a planned course of therapy is modified to include other treatment agents (alone or in combination) as a result of disease progression, relapse or toxicity, the exception to this being the need to attain a sufficient response for stem cell transplantation to proceed. A new line of therapy also starts when a planned period of observation off therapy is interrupted by a need for additional treatment for the disease.				
	Lenalidomide	The 3rd or later line of treatment in transplant ineligible patients with multiple	5. The patient is of ECOG performance status 0 or 1 or 2. Please tick one of the boxes below:	-			
LEN3	in combination with	myeloma previously treated with at least 2		No	TA171	18-Jun-09	16-Sep-09
	dexamethasone	prior regimens where the following criteria are met:	performance status 2				
		Citeria are met.	6. The patient has had no previous therapy with lenalidomide.				
			7. Lenalidomide is to be used in combination with either dexamethasone or dexamethasone plus cyclophosphamide and that it is not to be used in combination with any other agents unless accompanied by a separate and specific blueteq treatment criteria form. If cyclophosphamide is used in combination with lenalidomide and dexamethasone, the cyclophosphamide must be initiated with the first cycle of lenalidomide plus dexamethasone and not as a result of disease progression whilst on lenalidomide and dexamethasone.	-			
			8. Lenalidomide plus dexamethasone is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment, whichever is the sooner.				
			9. A formal medical review as to whether treatment with lenalidomide in combination with dexamethasone continues or not will be scheduled to occur at least by the end of the first 8 weeks of treatment.				
			10. No treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve).				
			11. Lenalidomide will be otherwise used as set out in its Summary of Product Characteristics.				
			1. This application is made by and the first cycle of systemic anti-cancer therapy with lenalidomide will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			2. The patient has a confirmed diagnosis of transfusion-dependent anaemia due to low- or intermediate-1-risk myelodysplastic syndromes (MDS) associated with an isolated deletion 5q cytogenetic abnormality	-			
			3. The other therapeutic options (e.g. best supportive care including regular red blood cell transfusions) are insufficient or inadequate.				
			4. When starting lenalidomide the ANC is greater than (>) 0.5 x 10^9/L and/or platelet counts greater than (>) 25 x 10^9/L.				
			5. The patient is of ECOG performance status 0 or 1 or 2. Please tick one of the boxes below:	-			
		The treatment of myelodysplastic syndromes associated with an isolated	- performance status 0 or - performance status 1 or				
LEN4	Lenalidomide	deletion 5q cytogenetic abnormality	performance status 2	No	TA322	24-Sep-14	23-Dec-14
		where the following criteria are met:	6. The patient has had no previous therapy with lenalidomide.				
			7. Lenalidomide is only to be used as a single agent at a starting dose of 10mg daily as per the summary of product characteristics	1			
			8. Lenalidomide is to be discontinued if no response after 4 cycles. If patients are responding after 4 cycles, lenalidomide will be continued until loss of response (progression of MDS or need for RBC transfusion) or unacceptable toxicity or patient choice to stop treatment, whichever is the sooner.	-			
			9. A formal medical review as to whether treatment with lenalidomide continues or not will be scheduled to occur at least by the end of the first 4 cycles of treatment.	1			
			10. No treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve).	1			
			11. Lenalidomide will be otherwise used as set out in its Summary of Product Characteristics.	1			

llueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is made by and the first cycle of systemic anti-cancer therapy with lenalidomide in combination with rituximab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient is an adult and has a histological diagnosis of follicular lymphoma of grades 1-3.  3. The patient has been previously treated with at least 1 prior systemic therapy for follicular lymphoma and now requires further systemic treatment.  For patients who have received rituximab or obinuturumab, please mark below as to whether the patient has disease that is anti-CD20 antibody sensitive i.e. responded to the last anti-CD20 antib				
LENS	Lenalidomide in combination with rituximab	For previously treated follicular lymphoma (grades 1-3a) where all the following criteria have been met:	Anti-CD20 antibody-resistant i.e. failed to respond to the last anti-CD20 antibody-containing regimen or had progressive disease within 6 months of completion of that anti-CD20 antibody-containing regimen  4. The patient is of ECOG performance status 0 or 1 or 2.  5. The patient has had no previous therapy with lenalidomide.  6. The patient will be treated with a maximum of 12 4-weekly cycles of lenalidomide.  7. The rituximab schedule of administration of 375mg/m2 given intravenously (IV) on days 1, 8, 15 and 22 in cycle 1 and then either 375mg/m2 given intravenously (IV) or 1400mg given subcutaneously (SC) on D1 only in cycles 2-5 will be used  8. Lenalidomide is only to be used in combination with rituximab and that it is not to be used in combination with any other agents.  Note: if rituximab has to be discontinued for toxicity, lenalidomide can be continued up to the maximum of 12 cycles.  9. Prior to cycle 1 the patient will receive tumour lysis syndrome prophylaxis (allopurinol, rasburicase or equivalent as per institutional guideline) and that the patient will be counselled as to be well orally hydrated during the 1st week of the 1st cycle or longer if clinically indicated.  10. The patient will have routine blochemistry tests performed weekly during cycle 1 and as clinically indicated and these results will be reviewed on day of testing to check for tumour lysis syndrome and its consequences.	No	TA627	07-Apr-20	06-Jul-20
			12. The guested with or severated on any formount have resections agreed on the severation of the semantion				
LEN6_v1.3	Lenalidomide	Lenalidomide monotherapy as maintenance treatment in newly diagnosed patients with multiple myeloma who have undergone autologeous stem cell transplantation where the following criteria have been met:	1. This application for maintenance lenalidomide is being made by and the first cycle of systemic anti-cancer therapy with maintenance lenalidomide monotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has newly diagnosed multiple myeloma.  3. The patient has recently undergone autologous stem cell transplantation.  4. The patient has had an adequate hemantological recovery following autologous stem cell transplantation.  5. Just prior to this application the patient has been tested for and has no evidence of disease progression since the transplantation was done.  6. The prescribing clinician understands that maintenance lenalidomide is recommended to start at about day 100 after stem cell transplantation.  7. The patient has been between the mumber of days since stem cell transplantation:  7. The patient has been previous therapy with lenalidomide unless the patient has been previously treated with 1st line lenalidomide allowed for transplant eligible patients via the interim treatment change options available during the coronavirus pandemic (blueted form LENIALCY will previously have been completed) or if the patient has been receiving NHS approved free of charge supply of maintenance lenalidomide as part of the NIHR RADAR trial and whilst still in remission has chosen to exit the trial or the patient has been receiving NHS approved free of charge supply of maintenance lenalidomide as part of the NIHR RADAR trial and whilst still in remission has chosen to exit the trial or the patient has been receiving NHS approved free of charge supply of maintenance lenalidomide as part of the NIHR RADAR trial and whilst still in remission has chosen to exit the trial or a fertile patient has been previously treated with 1st line lenalidomide (only in combination with dexamethasone) allowed for transplant eligible patients via the interim cancer treatment options available during the coronavirus pandemic (blueted form LENI	No	TA680	03-Mar-21	01-Jun-21
			8. The patient has an ECOG performance status of 0 or 1 or 2.  9. The patient will start maintenance lenalidomide at a dosing schedule of 10mg daily given on days 1:21 of a 28-day cycle and that any dose delays and reductions will be according to the Myeloma XI protocol version 9.0 (dated 2 November 2017).  Note: this dosing schedule is not the licensed one as set out in the lenalidomide Summary of Product Characteristics but is the one on which NICE assessed the clinical and cost effectiveness of maintenance lenalidomide. Note: the licensed dosing schedule of maintenance lenalidomide is not to be used.  10. My hospital Trust's governance policy regarding the use of unlicensed treatments has been followed as I understand that the above Myeloma XI dosing schedule of maintenance lenalidomide is unlicensed.  11. Lenalidomide is only to be used as monotherapy and that it is not to be used in combination with any other agents.  12. Lenalidomide is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment, whichever is the sooner.  13. A first formal medical review as to whether treatment with maintenance lenalidomide monotherapy continues or not will be scheduled to occur at least by the end of the first 8 weeks of treatment.  14. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment.				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
LNV1	Lenvatinib with everolimus	The treatment of previously treated advanced renal cell carcinoma	1. The application has been made by and the first cycle of systemic anti-cancer therapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The patient has a confirmed histological diagnosis of renal cell carcinoma with a clear cell component Note: papillary, chromophobe and Xp11 translocation sub types can be treated as per clear cell pathway  3. The patient has either metastatic disease or inoperable locally advanced disease  4. The patient has previously received only 1 vascular endothelial growth factor (VEGF)-targeted systemic therapy for advanced/metastatic renal cancer*  5. The patient has progressed on previous treatment or within 6 months of discontinuing previous treatment  6. The patient has an ECOG performance status of either 0 or 1*  *Patients with a performance status of 2 or more are not eligible for lenvatinib with everolimus  7. The patient has received no previous treatment with either lenvatinib or everolimus  8. The patient has no brain metastases or, if the patient has brain metastases, then these have been treated and are symptomatically stable  9. Lenvatinib with everolimus will be continued until loss of clinical benefit or unacceptable toxicity or patient choice to stop treatment  10. If unacceptable toxicity occurs, the daily doses of lenvatinib and, if necessary, everolimus are to be modified as needed according to the dose/management plan as set out in section 4.2 of the Summary of Product Characteristics for lenvatinib (Kisplyx)  11. No treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve)  12.Lenvatinib (Kisplyx) and everolimus are to be otherwise used as set out in their Summaries of Product Characteristics	No	TA498	24-Jan-18	24-Apr-18
LNV2	Lenvatinib	The treatment of differentiated thyroid cancer after radioactive iodine where all the following criteria are met:	1. This application is made by and the first cycle of systemic anti-cancer therapy with lenvatinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. This patient has a confirmed histological diagnosis of differentiated thyroid carcinoma (papillary or follicular or Hurtle cell type)  3. The patient has either metastatic disease or inoperable locally advanced disease  4. The disease is refractory to radioactive lodine  5. The disease is progressive and is either symptomatic or imminently likely to become symptomatic  6. The patient is treatment naive to both lenvatinib and sorafenib unless either: a) previously enrolled in the company's lenvatinib company's lenvatinib company secretary and the conditions set out in it) below or b) the patient has intolerant of sorafenib according to the conditions set out in it) below or b) the patient has had to discontinue sorafenib within 3 months of starting sorafenib because of toxicity (let there is sorafenib toxicity which cannot be managed by dose delay or dose modification) and there has been no disease progression whilst on sorafenib  Note: Sequential use of lenvatinib and then sorafenib is only funded if the patient has to discontinue lenvatinib because of intolerance within 3 months of its start and if the disease has not progressed whilst the patient is on lenvatinib. The use of lenvatinib after disease progression on or after sorafenib is not funded and vice versa.  7. The patient has an ECOG performance status of 0 or 1 or 2  8. Lenvatinib is to be continued as long as clinical benefit is observed or until there is unacceptable toxicity or patient choice to stop treatment  10. No treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve)  11. Lenvatinib is to be otherwise used as set out in its Summany of Product Characteristics	No	TA535	08-Aug-18	06-Nov-18

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
LNV3	Lenvatinib monotherapy	Treatment of Child-Pugh A locally advanced or metastatic hepatocellular carcinoma where the following criteria are met:	1. This application has been made by and the first cycle of systemic anti-cancer therapy with lenvatinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. One of the following applies to the patient, either:  - option 1 in which the patient has a confirmed histological diagnosis of hepatocellular carcinoma (HCC) or - option 2 in which a bloopy is deemed to be very high risk or technically not feasible in the patient and the criteria below are also all met:  a. the decision not to biosy has been made and documented by a specialist HCC multi-disciplinary team meeting  b. the tumour meets the non-invasive diagnostic criteria of HCC?  c. data is submitted as part of the ongoing "Systemic Therapy Audit, previously known as the Sorrafenib Audit 2.  It is expected that option 2 will only apply in exceptional circumstances and it should be noted that audit of non-biopsy rates will be reviewed regularly.  *EASL-CROTC Clinical Practice Guidelines: Management, Journal of Hepatology 2012 vol 56 p908-943. Non-invasive criteria can only be applied to cirrbotic patients and are based on imaging techniques obtained by 4-phase multidetector CT scan or dynamic contrast-enhanced MRI. Diagnosis should be based on the identification of the typical halimank of HCC (hypervascular in the arterial phase with washout in the portal venous or delayed phases). While one imaging technique is required for nodules beyond Izm in diameter, a more conservative approach with 2 techniques is recommended in suboptimal settings.  3. The patient has either metastatic disease or locally advanced disease that is ineligible for or failed surgical or loco-regional therapies  4. Either:  - the patient has not received any previous systemic therapy for hepatocellular carcinoma (option 1) or  - the patient has not received any previous systemic therapy for hepatocellular carcinoma (option 1) or  - the patient has not received any previous systemic therapy for hepatocellular carc	No	TA551	19-Dec-18	19-Mar-19

lueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
lueteq Form ref:	Lenvatinib in combination with pembrolizumab	Lenvatinib in combination with pembrolizumab for use in treatment-naive patients with intermediate or poor risk advanced renal cell carcinoma for whom treatment with nivolumab plus ipilimumab would otherwise be suitable where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy.  2. The precipitation of the provided provided in the control of the provided provid	drug/ indication	TASS8	NICE	baseline funding
			11. A formal medical review to assess the tolerability of treatment with lenvatinib plus pembrolizumab will be scheduled to occur at least by the start of the 3rd 3-weekly cycle or 2nd 6-weekly cycle of treatment and thereafter on a regular basis.  12. Treatment breaks of up to 12 weeks beyond the expected 3- or 6-weekly cycle length are allowed but solely to allow any toxicities to settle.  13. If the disease progresses on the pembrolizumab and lenvatinib combination and further systemic therapy is appropriate, the next line of treatment will be chosen from those options which are routinely commissioned ie for the next line of systemic therapy, there will be use of one choice of the following (mainly incorporating TKI options which have multiple modes of action): the currently commissioned 2nd line options of cabozantinib or everolimus monotherapy or the currently commissioned 1st line options of sunitinib (still on label as 2nd line treatment) or pazopanib (off label as 2nd line treatment) or tivozanib (off label as 2nd line treatment).  14. Lenvatinib and pembrolizumab will be otherwise prescribed and administered as outlined in their respective Summary of Product Characteristics (SPCs).				

v1.379 26-Nov-2025

Blueteq Form re	: Drug N	IICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
LIS01a	treat large gr met folli patie  chem other cell tr to 1: yet poten the poten the cells yet foll gr in fol gr in foll gr i	isocabtagene maraleucel for the timent of relapsed/refractory diffuse to 8-cell lymphoma (DLBCL) or high rade B-cell lymphoma or primary diastinal large B-cell lymphoma or cular lymphoma grade 38 either in ents who relapsed within 12 months of completion of 1st line noimmunotherapy AND who would wise be intended for potential stem ransplantation or who are refractory st line chemoimmunotherapy AND would otherwise be intended for this stem cell transplantation where to following criteria have been met:  This form is for the approval of pheresis and manufacture of CAR-T. There is a second part to this form helates to the subsequent infusion AR-T cells and this will be available er submission of the first part. The dpart of the form (US1a) and must be mart of the form (US1a) and must be mpleted on infusion of CAR-T cells submission of the first part of the form (US1a) and must be mpleted on infusion of CAR-T cells submission of the form (US1a) and must be mpleted on infusion of CAR-T cells submission of the first part of the form (US1a) and must be mpleted on infusion of CAR-T cells submission of the first part of the form (US1a) and must be mpleted on infusion of CAR-T cells submission of the first part of the form (US1a) and must be mpleted on infusion of CAR-T cells submission of the first part of the form (US1a) and must be mpleted on infusion of CAR-T cells submission of the first part. The manufacture of the form (US1a) and must be made and the manufacture of the form (US1a) and must be made and the manufacture of the form (US1a) and must be made and the manufacture of the form (US1a) and must be made and manufacture of the form (US1a) and must be made and must be made and manufacture of the form (US1a) and must be made and manufacture of the form (US1a) and must be made and manufacture of the form (US1a) and must be made and manufacture of the form (US1a) and must be made and manufacture of the form (US1a) and must be made	1. This againstant is being made by an official or footblood processing of presentation of the search protection of contingent specifically raised and screentation in the search of private content of the search protection of the search of the search protection of the search of the search of the search protection of the search of the s	No	TA1048	26-Mar-25	24-Jun-25

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lueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			8. The patient has been previously treated with a regimen containing an anti-CD20 monoclonal antibody unless there is clear documentation of the determination of CD20 negative disease.				
			9. On the date that the patient was confirmed as having refractory or relapsed disease according to the above definitions, the patient had only received 1st line of therapy for the DLBCL or HGBCL or PMBCL or FL3B or TFL to DLBCL or other transformed conditions to DLBCL.				
			Note: in the case of patients who have transformed from a lymphoma or other condition to DLBCL, 1st line therapy refers to the treatment of the disease (e.g. TFL to DLBCL) once transformation has been documented.				
			Note: it is recognised that some patients at the time of the demonstration of refractory or relapsed disease have very rapidly progressive disease and thus have to commence urgent 2nd line treatment. It is therefore acceptable for patients to have received a maximum of 2 cycles of standard 2nd line chemotherapy regimens with one of the following regimens ('anticipatory bridging therapy'): R-GDP, R-GemCarbo, R-ESHAP, R-ICE, R-IVE, R-BendaPola and the Marietta protocol.				
			Please enter below whether the rate of disease progression as outlined above required urgent 2nd line salvage chemotherapy ('anticipatory bridging therapy') in this patient: - no urgent chemotherapy required prior to this application or				
			- a maximum of 2 cycles of one of the above standard salvage chemotherapy regimens have been given prior to this application on grounds of urgent need and all other treatment criteria on this form are fulfilled				
		Lisocabtagene maraleucel for the treatment of relapsed/refractory diffuse large B-cell lymphoma (DLBCL) or high grade B-cell	10. In the absence of the availability of lisocabtagene maraleucel for this 2nd line indication the patient would have been fit and intended for both standard 2nd line salvage chemotherapy (see note below) and potential stem cell transplantation.				
			Note: Second line treatment regimens which are appropriate include: R-GDP, R-GemCarbo, R-ESHAP, R-ICE, R-IVE, R-BendaPola and the Marietta protocol.				
		lymphoma or follicular lymphoma grade 3B	11. The patient has not previously been treated with an anti-CD19 antibody-drug conjugate.				
		either in patients who relapsed within 12 months of completion of 1st line chemoimmunotherapy AND who would	12. Whether the patient has active CNS involvement by the lymphoma or not and if present whether this is in addition to systemic disease progression or not.				
		otherwise be intended for potential stem cel	Please tick one of the boxes below:				
		transplantation or who are refractory to 1st	- currently no known CNS involvement or - currently has both active CNS and systemic disease or				
		otherwise be intended for potential stem cel					
LISO1a	Lisocabtagene maraleucel	transplantation where the following criteria have been met:	Note: patients with primary CNS lymphoma are not eligible for treatment with lisocabtagene maraleucel.	No	TA1048	26-Mar-25	24-Jun-25
		This form is for the approval of leucapheresis and manufacture of CAR-T cells. There is a	13. The patient has an ECOG performance score of 0 or 1. Please enter below as to the patient's current ECOG performance status (PS):	-			
			The ECOG performance status scale is as follows:				
			PS 0 The patient is fully active and able to carry on all pre-disease performance without restriction				
		part. The second part of the form (US1b) car	PS 1 The patient is restricted in physically strenuous activity but is ambulatory and able to carry out work of a light or sedentary nature eg light housework, office work PS 2 The patient is ambulatory and capable of all selfcare but unable to carry out any work activities and is up and about more than 50% of waking hours				
		only be completed as a continuation of this	1-3.2 The platient is ambiliation yand capabile of an Setticate but unable to Carry out any work activities and is to up and about more trian 50% of waking hours 8-3.3 The platient is capable of only limited self-tiera and is confined to be do or chair more than 50% of waking hours				
		completed on infusion of CAR-T cells	PS 4 The patient is completely disabled, cannot carry out any selfcare and is totally confined to bed or chair				
		otherwise the treating Trust will not be reimbursed for the cost of lisocabtagene	The patient currently has a performance status of either				
		maraleucel	- ECOG PS 0 or - ECOG PS 1				
			14. The patient has sufficient end organ function to tolerate treatment with CAR-T cell therapy.				
			15. The patient has either had no previous therapy with any genetically modified autologous or allogeneic T cell immunotherapy or the patient has been treated with doses of genetically modified autologous or allogeneic T cell immunotherapy within an abandoned dosing cohort in a first in human dose-escalation phase I clinical trial.				
			Please tick appropriate box as to which type of previous treatment the patient has had:				
			- No previous therapy with any genetically modified autologous or allogeneic T cell immunotherapy or - Previously treated with doses of genetically modified autologous or allogeneic T cell immunotherapy within an abandoned dosing cohort in a first in human dose-escalation phase I clinical trial				
			16. Prior to infusion 2 doses of tocilizumab are available for use in this patient in the event of the development of cytokine release syndrome.	1			
			17. Lisocabtagene maraleucel-modified CAR-T cell therapy will otherwise be used as set out in its Summary of Product Characteristics (SPC).	1			
			18. Approval for the use of lisocabtagene maraleucel has been formally given by the National DLBCL/HGBCL CAR-T cell Clinical Panel.	1			
			Please state date of approval				

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Blueteq Form ref:	: Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
LISO1b	Lisocabtagene maraleucel	Lisocabtagene maraleucel for treating relapsed/refractory diffuse large B-cell lymphoma (DIBCL) or primary mediastina large B-cell lymphoma (BCCL) or primary mediastina large B-cell lymphoma (PMBCL) or follicular lymphoma grade 38 (FL3B) and in adult patients either who relapse within 12 months of completion of 1st line chemoimmunotherapy AND with chemoimmunotherapy AND who would otherwise be intended for potential stem cell transplantation or who are refractory to 1st line chemoimmunotherapy AND who would otherwise be intended for potential stem cell transplantation where the following criteria have been met:  This second part of the form is to document the date of infusion of CAR-T cell therapy and for registration of this infusion with NHS England so that the treating Trust is reimbursed for the cost of lisocabtagene maraleucel. There is a first part of the form for the approval of leucapheresis and manufacture of CAR-T cells which has already been completed (LISLa). This second part of the form (LISLb) should only be completed as a continuation form once the date of CAR-T cell infusion is known.	- no bridging therapy at all or - corticosteroids only or - chemo(immuno)therapy only with intensive salvage-type therapy (eg R-GDP, R-GemCarbo, R-ESHAP, R-ICE, R-IVE, R-BendaPola and the Marietta protocol) or - chemo(immuno)therapy only with BR-polatuzumab or - other chemo(immuno)therapy only or - radiotherapy only or - corticosteroids and chemo(immuno)therapy or - corticosteroids and radiotherapy or - chemo(immuno)therapy and radiotherapy or - chemo(immuno)therapy and radiotherapy ± corticosteroids  4. The nature of any imaging procedure performed to assess response to bridging therapy below:  no bridging therapy and so no radiological assessment performed or - CT or MR scan performed or - CT or MR scan performed or	- No	TA1048	26-Mar-25	24-Mar-25

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. I confirm that this application is made by and the first cycle of systemic anti-cancer therapy with liposomal cytarabine and daunorubicin will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy				
		The treatment of adults with newly diagnosed acute myeloid leukaemia (AML)	2. This patient is an adult and has a confirmed diagnosis of acute myeloid leukaemia with one of the following types:  - therapy-related AML (t-AML) with a documented history of prior cytotoxic therapy or ionising radiotherapy for an unrelated disease or  - chronic myelomonocytic leukaemia AML (CMMoL AML) with a documented history of CMMoL prior to transformation to AML or  - myelodysplasia AML (MDS AML) with a documented history of MDS prior to transformation to AML or  - de novo AML with karyotypic changes characteristic of MDS.				
LCD1	Liposomal cytarabine and daunorubicin	that is secondary to therapy or myelodysplasia or chronic myelomonocytic leukaemia where the following criteria are	3. I confirm that the patient is newly diagnosed with one of the above types of AML and has not received any chemotherapy for this AML.  4. I confirm that the patient has an ECOG performance score of 0, 1 or 2.	No	TA552	19-Dec-18	19-Mar-19
		met:	** Tournitrials the patient is fifted induction chemotherapy with liposomal cytarabine and daunorubicin.				
			6. I confirm that the patient will be treated with liposomal cytarabine and daunorubicin with the doses and schedules for induction chemotherapy as outlined in the Summary of Product Characteristics of liposomal cytarabine and daunorubicin.				
			7. I note that the use of liposomal cytarabine and daunorubicin is exempt from the NHS England Treatment Break policy	1			
			8. I confirm that liposomal cytarabine and daunorubicin is to be otherwise used as set out in its Summary of Product Characteristics				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is being made by and the first cycle of systemic anti-cancer therapy.  2. The patient has a histologically confirmed diagnosis of diffuse large B cell lymphoma (DLBCL) or high grade B cell lymphoma or transformed follicular lymphoma to DLBCL.  The definition of DLBCL includes the following:  - DLBCL not otherwise specified (NOS) [including germinal centre B-cell (GCB) and activated B-cell (ABC) subtypes]  - primary mediastinal large B cell lymphoma  - T cell rich B cell lymphoma  - T cell rich B cell lymphoma  - T cell rich B cell lymphoma  - Ingit grade B-cell lymphoma (double hit and triple hit high grade B cell lymphoma)  Note: Patients with primary (CNS lymphoma, Burkitt lymphoma and plasmablastic lymphoma) are NOT eligible for treatment with loncastuximab tesirine.  Please record in the box below whether the patient has DLBCL according to one of the types within the above definition OR  - the patient has to BLBCL according to one of the types within the above definition OR  - the patient has DLBCL according to one of the types within the above definition OR  - the patient has DLBCL cording to one of the rypes within the above definition OR  - the patient has DLBCL or TFL to DLBCL either of which has relapsed following or during 2 or more lines of standard routinely commissioned systemic therapies and that within these 2 lines of therapy there has been treatment with an anti-CD20 regimen and an anti-cD20 regimen and an anti-according regimen.  Note: patients with TFL to DLBCL with the patient has received 2 or more lines of systemic therapy given specifically for the transformed follicular lymphoma to DLBCL  4. The number of lines of systemic therapy that the patient has received 2 or more lines of systemic therapy. Similarly, bridging chemotherapy prior to and then followed by CAR T cell therapy counts as 1 line of systemic therapy. Similarly, bridging chemotherapy prior to and then followed by CAR T cell therapy counts as 1 line of systemic therapy. Similarly, bridging chemotherapy pri				
LON1_v1.0	Loncastuximab tesirine monotherapy	For the further treatment of adult patients with diffuse large B-cell lymphoma or high grade B-cell lymphoma who have received previous treatment with 2 or more lines of systemic therapy (which have included polatuzumab vedotin unless the use of polatuzumab vedotin was contra-indicated) and in addition are not candidates for any future CAR T-cell therapy where the following criteria have been met:	5. The patient has been previously treated with stem cell transplantation:  - No previous stem cell transplantation OR  - Yes, previous stem cell transplantation  6. The patient has been previously treated with CAR T cell therapy and if so at which place in the treatment pathway:  - No previous CAR T cell therapy and the patient is unsuitable for CAR T cell therapy both now and in the future OR  - Yes, previous CAR T cell therapy as 2nd line therapy OR  - Yes, previous CAR T cell therapy as 3rd or more line of therapy  Note: Swedish Orphan Biovitrum (the company that markets loncastuximab tesirine) did not make an evidence submission to NICE for consideration of the use of loncastuximab tesirine in patients who are suitable for or might become eligible for CAR T cell therapy.  Note: Ioncastuximab tesirine must not be used as bridging therapy for CAR T cell therapy.  Note: Doncastuximab tesirine must not be used as bridging therapy for CAR T cell therapy must be re-biopsied prior to consideration of treatment with loncastuximab tesirine to ensure that the lymphoma retains CD19 protein expression (unless such a biopsy is unsafe for the patient and the patient has progressive disease at previously known sites of active disease and the previous histology was DLBCL or HGBCL).	No	TA947	31-Jan-24	30-Apr-24
			7. The patient has either previously received systemic therapy with a regimen containing polatuzumab vedotin or the use of a polatuzumab vedotin-containing regimen was contraindicated.  Note: the NICE recommendation for access to loncastuximab tesirine stipulates that for treating relapsed or refractory DLBC. In patients who have had 2 or more systemic therapies, lonastuximab is only recommended if patients have received prior polatuzumab (whether relapsed following such treatment or refractory to it or if polatuzumab was not tolerated) or if treatment with polatuzumab was contraindicated.  Please record in the box below which of the following applies to this patient:  - previous treatment with 1st line polatuzumab vedotin-containing chemotherapy to which the patient had relapsed or refractory disease OR  - previous treatment with 1st or greater line polatuzumab vedotin-containing chemotherapy to which the patient had relapsed or refractory disease OR  - previous treatment with a or greater line polatuzumab vedotin-containing chemotherapy which was not tolerated and hence treatment with a polatuzumab vedotin or containing chemotherapy which was not tolerated and hence treatment with polatuzumab vedotin for this reason  8. The patient has not been previously treated with loncastuximab tesirine unless loncastuximab tesirine has been accessed via a company compassionate access scheme and all other treatment criteria on this form are fulfilled.				
			9. The patient has an ECOG performance status score of 0 or 1 or 2.  10. Loncastuximab tesirine is to be administered as monotherapy and not in combination with any other systemic therapies for lymphoma.  11. The dosing schedule of loncastuximab tesirine differs in cycle 3 and beyond from that used in cycles 1 and 2.  12. Treatment with loncastuximab tesirine monotherapy will be stopped at whichever of the following events occurs first: disease progression or unacceptable toxicity or withdrawal of patient consent.  Note: there is no formal stopping rule for loncastuximab tesirine in this indication but once loncastuximab is electively stopped (ie not for reasons of toxicity), it cannot be re-started.  13. The prescribing clinician and the treating team are familiar with the dose modifications and delays required for the management of adverse reactions to loncastuximab tesirine, both haematological and non-haematological (eg for oedema, effusions, cutaneous toxicity and abnormal liver function tests).				
			14. A formal medical review as to whether treatment with loncastuximab tesirine should continue or not will be scheduled to occur at least by the end of the first 6 weeks of treatment.  15. When a treatment break of more than 6 weeks beyond the expected 3-weekly cycle length is needed, a treatment break approval form will be completed to restart treatment  16. Loncastuximab tesirine will be otherwise used as set out in its Summary of Product Characteristics (SPC)				

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application for lorlatinib is being made by and the first cycle of systemic anti-cancer therapy with lorlatinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			2. The patient has a locally advanced or metastatic non-small cell lung cancer.				
			3. The patient has a histologically or cytologically confirmed diagnosis of non-small cell lung cancer that carries an anaplastic lymphoma kinase (ALK) rearrangement based on a validated test.				
LORI	Lorlatinib	For anaplastic lymphoma kinase positive advanced non-small-cell lung cancer previously treated with 1st line alectinib or 1st line brigatinib or 1st line brigatinib or 1st line brigatinib or 1st line brigatinib or 1st line certitinib or 1st tyrosine kinase inhibitor therapy (brigatinib tyrosine kinase inhibitor therapy (brigatinib or certitinib) or after disease progression during adjuvant alectinib or within 6 months of completion of adjuvant alectinib where 6 the following criteria have been met:	4. The only TXI treatment that the patient has progressed on is 1st line alectinib or 1st line critoriin bro 1st line patient has progressed on:  - 1st line lactinib or - 1st line certinib bro 1st line certinib bro 1st line certinib bro 1st line critoriin bro 1st line critor	No	TA628	13-Мау-20	11-Aug-20
			11. A formal medical review as to whether treatment with lorlatinib should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment.  12. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment.				
			13. Lordatinib will be otherwise used as set out in its Summary of Product Characteristics.  1. This application is made by a consultant oncologist who is specifically trained and accredited in the use of systemic anti-cancer therapy and who is a core member of the relevant Neuroendocrine Carcinoma Multi-Disciplinary Team (MOT)			+	
			2. The Neuroendocrine Carcinoma MDT has confirmed the arrangements by which only persons authorised to handle radiopharmaceuticals (such as Jutetium oxodotreotide) do so in authorised clinical settings and after evaluation of the patient by an appropriately trained and accredited physician				
			3. The patient has a histologically documented, well differentiated neuroendocrine carcinoma of the gastrointestinal tract or pancreas  Note: patients with primary bronchial neuroendocrine carcinomas are not eligible for treatment with lutetium oxodotreotide				
		Lutetium oxodotreotide for unresectable	4. The patient's disease is either unresectable or metastatic				
		or metastatic, progressive, well	5. The patient's disease is somatostatin receptor positive on imaging (on PET scanning but otherwise on scintigraphy if PET scanning not possible) and this imaging confirms overexpression of somatostatin receptors in the tumour tituse with the fundor under the least as high as normal liver under keep rade core 2 control tituse with the fundor under the least as high as normal liver under keep rade core 2 control tituse with the fundor under the least as high as normal liver under keep rade core 2 control to the least as high as normal liver under keep rade core 2 control tituse with the least as high as normal liver under keep rade core 2 control tituse with the least as high as normal liver under keep rade core 2 control tituse with the least as high as normal liver under keep rade core 2 control tituse with the least as high as normal liver under keep rade core 2 control tituse with the last as high as normal liver under keep rade core 2 control tituse with the last as high as normal liver under keep rade core 2 control tituse with the liver as a li				
LUT1	Lutetium oxodotreotide	differentiated and somatostatin receptor	Issue with the tunior upoke at less as high as home mer upoke (tunior upoke) and (for MR imaging over the course of a maximum period of 3 years  6. There has been exent demonstration in this patient of disease progression on CF or MR imaging over the course of a maximum period of 3 years	No	TA539	29-Aug-18	27-Nov-18
10.1	Lutetium oxodoti eotide	positive gastroenteropancreatic			12333	25-Aug-10	27-1404-10
		neuroendocrine carcinoma where all the following criteria are met:	7. The patient has an ECOG performance status (PS) score of 0 or 1 or 2				
		Tollowing criteria are met.	8. The patient has not received prior treatment with lutetium oxodotreotide Note: re-treatment with a further program of lutetium oxodotreotide treatments is not commissioned				
			9. Lutetium oxodotreotide is being given as monotherapy (bar somatostatin analogues in between treatments) and will involve a maximum of 4 infusions of 7400 MBq as long as there is no evidence of disease progression				
			10. A formal face to face medical review as to whether treatment with lutetium oxodotreotide should continue or not will be scheduled to occur before each of the 4 planned treatment administrations				
		11. Tr	11. The presciribing clinician notes that the use of lutetium oxodotreotide is exempt from the NHS England cancer drug Treatment Breaks policy				
			12. Lutetium oxodotreotide will otherwise be used as set out in its Summary of Product Characteristics (SPC)				

Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
Midostaurin	Midostaurin for treating newly diagnosed FLT3 mutation positive acute myeloid leukaemia (FLT3-TIO or FLT3-TIO) in ADULTS where the following criteria are met:	1. An application is being made by and the first cycle of systemic anti-cancer therapy with midostaurin will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. This patient is an adult and has a confirmed diagnosis of acute myeloid leukaemia  3. The patient's AML has a FLT3 mutation (ITD or TKD) as determined by a validated test:  Please mark below which type of FLT3 mutation applies to this patient:  - ITD disease or  - TKD disease  - TKD disease or  - TKD disease or  - TKD disease  - TKD disease	No	TAS23	13-Jun-18	11-Sep-18
		Note: the use of midostaurin after a stem cell transplant is not commissioned.  10. Midostaurin is to be otherwise used as set out in its Summary of Product Characteristics  1. This application for midostaurin monotherapy is being made by and the first cycle of systemic anti-cancer therapy with midostaurin monotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a pathologically-confirmed diagnosis of aggressive systemic mastocytosis (ASM) or aggressive systemic mastocytosis with an associated haematological neoplasm (ASM-AHN) or mast cell leukaemia.				
Midostaurin	For aggressive systemic mastocytosis or aggressive systemic mastocytosis with an associated haematological neoplasm or mast cell leukaemia where the following criteria have been met:	- aggressive systemic mastocytosis (ASM) - aggressive systemic mastocytosis with an associated haematological neoplasm (ASM-AHN) - mast cell leukaemia  3. The patient has advanced disease and requires systemic therapy for this condition.  4. Either the patient has received previous systemic therapy for this condition or not.  Please mark below whether the patient has/has not previously received any systemic therapy for this condition: - no, this patient has not received any previous systemic therapy for this condition - yes, this patient has been previously treated with systemic therapy for this condition  5. Either the patient has received previous treatment with avapritinib or not.  Please mark below whether the patient has previously received avapritinib - yes, the patient has not received any previous avapritinib - yes, the patient has been previously treated with avapritinib - yes, the patient has been previously treated with avapritinib - yes, the patient has not previously received treatment with middsaurin.  Note: If natients were entered into the company's early argests/compansionate use scheme for middstaurin for these indications they must continue to greate middstaurin from this scheme. These natients must not be transferred to	No	TA728	22-Sep-21	21-Dec-21
	Midostaurin	Midostaurin Midostaurin Midostaurin Midostaurin  ADULTS where the following criteria are met:  For aggressive systemic mastocytosis or aggressive systemic mastocytosis or aggressive systemic mastocytosis with an associated haematological neoplasm or mast cell eliekaemia where the following criteria celle with the mastocytosis with an associated haematological neoplasm or mast celle with the mastocytosis with an associated haematological neoplasm or mast cell eliekaemia where the following criteria.	An equipation is bring raise by and finding point grown anticoners through an industration will be prevailed by a consultant special by special price and a construction of the large of the price of	Process   Proc	Part   Part	Manual Part

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
MID3	Midostaurin	For treating newly diagnosed FLT3 mutation positive acute myeloid leukaemia (FLT3-ITD or FLT3-TKD) in POST PUBESCENT CHILDREN LESS THAN 18 YEARS OLD where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with midostaurin will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient is a post pubsecent child less than 18 years old and has a confirmed diagnosis of acute myeloid leukaemia.  Note: midostaurin is not licensed for AML in this age group and hence completion of this form also confirms that Trust policy is being followed as regards the use of unlicensed medicines.  Note: For adults there is a separate blueten form.  3. The patient's AML has a FLT3 mutation (ITD or TKD) as determined by a validated test.  Please mark below which type of FLT3 mutation applies to this patient:  -ITD disease or  -ITD diseas	No	TA523	13-Jun-18	03-Feb-23

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
MOG1	Mogamulizumab	Mogamulizumab as 3rd line systemic therapy or beyond 3rd line systemic therapy for patients with stage IIB to Ne mycosis fungoides where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with mogamulizumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for adverse reactions to mogamulizumab and the prescribing clinician understands the need for testing for hepatitis B before mogamulizumab treatment commences and the risk of tumour lysis syndrome in patients with rapidly proliferating disease and high tumour burden.  3. The patient has a diagnosis of mycosis fungoides.  Please note that there is a separate form MOG2 for patients with Sezary syndrome.  4. The disease stage of mycosis fungoides is stage IIB to IVB.  Please mark below the stage of disease that applies to this patient:  - stage IIB mycosis fungoides  - stage IIA mycosis fungoides  - stage IVAI mycosis fungoides  - s		TA754	Guidance 15-Dec-21	_
			12. Mogamulizumab will be stopped at whichever of the following events occurs first: disease progression or unacceptable toxicity or withdrawal of patient consent.  13. A formal medical review as to how mogamulizumab is being tolerated and whether mogamulizumab should continue or not will be scheduled to occur at least by the end of the second month of treatment.  14. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, the prescribing clinician will complete a treatment break approval form to restart treatment, including as appropriate if the patient has had an extended break on account of Covid-19.  15. Mogamulizumab will otherwise be used as set out in its Summary of Product Characteristics (SPC) with the exception of criteria 4 and 5 above.				

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
MOG2	Mogamulizumab	Mogamulizumab as 2nd line systemic therapy or beyond 2nd line systemic therapy for patients with stage I/N to I/N Sezary syndrome where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy, with mogamulizumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for adverse reactions to mogamulizumab and the prescribing clinician understands the need for testing for hepatitist is before mogamulizumab treatment commences and the risk of tumour lysis syndrome in patients with rapidly proliferating disease and high tumour burden.  3. The patient has a diagnosis of Sezary syndrome.  Please meant below the stage of disease that applies to this patient: - stage NAI Sezary syndrome  9. The patient has received at least 1 line of systemic treatment for Sezary syndrome.  Note: mogamulizumab is only recommended by NCC if the patient has received at least 1 line of systemic therapy.  6. The patient has received at line systemic therapy for Sezary syndrome.  Note: mogamulizumab is only recommended by NCC if the patient has received at least 1 line of systemic therapy.  6. The patient has received at line systemic therapy for Sezary syndrome.  Note: mogamulizumab is only recommended by NCC if the patient has received at least 1 line of systemic therapy.  6. The patient has received at line systemic therapy was received by the patient: - ineterform	No	TA754	15-Dec-21	15-Mar-22

1. This application is being made by and the first cycle of systemic anti-cancer therapy.  2. The patient is an adult with a diagnosis of primary myelofibrosis (also known as chronic idiopathic myelofibrosis) or post polycythaemia vera myelofibrosis or post essential thrombocythaemia myelofibrosis.  Please enter below as to which type of myelofibrosis applies to this patient: - primary myelofibrosis - post polycythaemia vera myelofibrosis or - post polycythaemia vera myelofibrosis - post polycythaemia vera myelofibrosis or - post polycythaemia vera myelofibrosis - primary myelofibrosis - post polycythaemia vera myelofibrosis or post essential thrombocythaemia myelofibrosis - post polycythaemia vera myelofibrosis or post essential thrombocythaemia myelofibrosis - post polycythaemia vera myelofibr	Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
10. Momelotinib is to be continued as long as the benefit-risk remains positive for the patient.  11. The prescribing clinician is aware that momelotinib has clinically important interactions with various drugs which can affect the CYP3A4 and other enzyme systems and also transporters (as set out in sections 4.4 and 4.5 of 12. The prescribing clinician is aware of the risks of infection including Hepatitis B reactivation that can occur during treatment with momelotinib.  13. A formal medical review as to how momelotinib is being tolerated and whether treatment with momelotinib should continue or not will be scheduled to occur at least by the start of the third 4-weekly cycle of treatment.  14. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, the prescribing clinician will complete a treatment break approval form to restart treatment.	мом1	Momelotinib monotherapy	anaemic patients with myelofibrosis and disease-related splenomegaly or symptoms	2. The patient is an adult with a diagnosis of primary myelofibrosis (also known as chronic idiopathic myelofibrosis) or post polycythaemia vera myelofibrosis or post essential thrombocythaemia myelofibrosis.  Please enter below as to which type of myelofibrosis applies to this patient: - primary myelofibrosis or - post polycythaemia vera myelofibrosis or - post polycythaemia vera myelofibrosis or - post essential thrombocythaemia myelofibrosis  3. The patient's myelofibrosis has a risk category that is either intermediate-2 or high risk.  Please enter below which myelofibrosis risk category applies to this patient: - intermediate-2 risk or - high risk - The patient has disease-related splenomegaly or symptoms.  5. The patient has moderate to severe anaemia. 6. The patient has been previously treated with rusolitinib or not no previous treatment with rusolitinib or or - previous treatment with rusolitinib or - yes, the patient has been previously treated with rusolitinib - yes, the patient has been previously treated with rusolitinib - yes, the patient has been previously treated with rusolitinib - yes, the patient has been previously treated with rusolitinib - yes, the patient has been previously treated with rusolitinib - yes, the patient has been previously treated with rusolitinib - yes, the patient has been previously treated with rusolitinib - yes, the patient has been previously treated with rusolitinib - yes, the patient has been previously treated with rusolitinib - yes, the patient has been previously treated with rusolitinib - yes, the patient has been previously treated with rusolitinib - 1. The patient has not Pock patient with rusolitinib - 2. The patient has been previously treated with rusolitinib - 3. The patient has been previously treated with rusolitinib - 3. The prescribing clinical is as ware the patient has been previously treated with rusolitinib - 3. The prescribing clinical is as ware that momelotinib has clinically important interactions with various drugs which can affect the CYP3	No	TA957	20-Mar-24	18-Jun-24

ueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. An application has been made by and the first cycle of systemic anti-cancer therapy with nab-paclitaxel will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy				
			2. The patient has a confirmed histological or cytological diagnosis of breast cancer.			I	ĺ
			3. The patient is being switched to nab-paclitaxel from either paclitaxel or docetaxel either following a severe hypersensitivity reaction which precludes further exposure to paclitaxel or docetaxel or to reduce the risks of treatment in potentially vulnerable patients				
NAB1	Nab-Paclitaxel	Paclitaxel as albumin-bound nanoparticles (nab-paclitaxel) for breast cancer where the following criteria have been met:	4. Nab-paclitaxel is to be used either as a single agent or in combination for  - neoadjuvant treatment - adjuvant treatment - treatment of metastatic disease	No			
			5. The licensed dose of nab-paclitaxel at 260mg/m2 IV every 21 days will be used when given as monotherapy.				Í
			Note: The dose may be attenuated when given in combination with other chemotherapies.  6. The patient has an ECOG performance status of 0, 1 or 2.				İ
			7. Trust policy regarding the use of unlicensed treatments has been followed as nab-paclitaxel is not licensed for use in early breast cancer. (It is only licensed for use in metastatic breast cancer)				
			8. Nab-paclitaxel will otherwise be used as set out in its Summary of Product Characteristics (SPC).	1		<u> </u>	1
			1. This application is being been made by and the first cycle of systemic anti-cancer therapy with nab-paclitaxel plus gemcitabine will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.	i.		06-Sep-17	
			2. The patient has confirmed histological or cytological diagnosis of pancreatic adenocarcinoma.				1
			3. The patient has metastatic disease (patients with locally advanced disease are ineligible).				İ
NAB2	Nab-paclitaxel with gemcitabine	The treatment of untreated metastatic pancreatic cancer only if other combination chemotherapies are unsuitable and they would otherwise have gemcitabine monotherapy	4. The patient is either completely treatment naïve for systemic therapy for pancreatic cancer or the patient has received prior systemic anti-cancer therapy as neo-adjuvant or adjuvant therapy AND such treatment was completed at least 6 months previously.  Please mark below whether or not previous systemic anti-cancer therapy for pancreatic cancer has ever been received in the neoadjuvant or the adjuvant disease settings:  - no previous neoadjuvant/adjuvant systemic therapy of any kind and treatment naïve for metastatic pancreatic cancer  - prior neoadjuvant chemotherapy for non-metastatic disease and the last dose received by the patient was 6 or more months prior to this application  - prior chemotherapy in the adjuvant setting and the last dose received by the patient was 6 or more months prior to this application	No	TA476		05-Dec-17
		generaline monocherapy	5. Nab-pacilitaxel is to be used only in combination with gemcitabine.				ĺ
			6. Nab-pacilitaxel plus gemcitabine is to be used as 1 <sup>st</sup> line treatment only.				1
			7. The patient has a performance status of 0 or 1.	1			1
			8. The patient is not considered to be a suitable candidate for oxaliplatin- and irinotecan-based combination chemotherapy and would otherwise receive gemcitabine monotherapy.				ĺ
			9. Nab-paclitaxel will otherwise be used as set out in its Summary of Product Characteristics (SPC).				İ
		The treatment of refractory T-cell acute lymphoblastic leukaemia or refractory T-	1. Application made by and first cycle of systemic anti-cancer therapy to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy		-/- NUIC Fardand		
NEL1	Nelarabine	cell lymphoblastic non-Hodgkin's	2. a) Refractory T-cell acute lymphoblastic leukaemia, OR	Yes	n/a - NHS England clinical policy	-	01-Apr-21
		lymphoma where all the following criteria are met:	b) Refractory T-cell lymphoblastic non-Hodgkin's lymphoma	1			ĺ
		are met.	3. Treatment intent is to proceed to bone marrow transplantation	1	1		į.

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
NER1	Neratinib	The extended adjuvant therapy for hormon receptor positive HER2-overexpressed early breast cancer after completion of adjuvant therapy with HER2 targeted monotherapy with trastuzumab where the following criteria have been met:	1. This application for neratinib as extended adjuvant chemotherapy is made by and the first cycle of adjuvant neratinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has histologically documented breast cancer which is BOTH hormone receptor positive and HER2 overexpressed (HER2 3+ by immunohistochemistry and/or has a ratio of 22.0 by in situ hybridisation). Note: neratinib is not licensed for extended adjuvant therapy in hormone receptor negative patients.  3. The patient has been diagnosed with early breast cancer and this has been adequately excised.  4. That either the patient did not receive necadjuvant therapy or the patient was treated with necadjuvant therapy AND there was residual invasive carcinoma in the breast and/or the axilla. Please mark below which applies to this patient:  - patient did not receive necadjuvant therapy or - patient did receive necadjuvant therapy or - patient did receive necadjuvant therapy and therapy and therapy resulted in a pathological complete remission or if there was only residual carcinoma in situ disease in the breast and a pathological complete remission in the axillary nodes (if the axillary hymph node status was positive prior to necadjuvant treatment).  5. The patient has received chemotherapy in the management of the early breast cancer either as necadjuvant treatment pre-definitive surgery or as adjuvant therapy post-surgery.  6. The patient has completed adjuvant therapy with treaturumab as HER2-targeted monotherapy and is within 1 year of completing such trastuzumab monotherapy.  Note: NICE has not recommended use of neratinib if the patient received any pertuzumab as part of adjuvant therapy. Patients treated with necadjuvant chemotherapy in combination with pertuzumab and trastuzumab are only eligible for neratinib therapy if the pertuzumab was solely used as part of necadjuvant treatment and no pertuzumab was used as part of adjuvant therapy.  7. The patient has a	No	TA612	20-Nov-19	18-Feb-20
N/A	Nilotinib	Nilotinib for the treatment of untreated chronic phase chronic myeloid leukaemia	12. Neratinib will be otherwise used as set out in its Summary of Product Characteristics (SPC)  1. I confirm that an application has been made and the first cycle of systemic anti-cancer therapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. I confirm that the patient has chronic phase myeloid leukaemia 3. I confirm that the patient has received no prior treatment 4. I confirm that intantinib is not appropriate for this patient and that this has been discussed and supported by the relevant MDT involved in CML decision making  5. I confirm that nilotinib will be used as outlined in the Summary of Product Characteristics (SPC).	No No	TA426	21-Dec-16	21-Mar-17
NIL4	Nilotinib	For treating imatinib-resistant or imatinib- intolerant Philadelphia chromosome positive chronic phase chronic myeloid leukaemia in children where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with nilotinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has Philadelphia chromosome positive CML in chronic phase.  3. The patient has been previously treated with imatinib which had to be discontinued due to resistance or intolerance.  Please mark below whether the patient was resistant to or intolerant of imatinib: - intolerant of imatinib  4. The use of nilotinib has been discussed by the relevant multi-disciplinary team (MDT) involved in chronic myeloid leukaemia (CML) decision making, which must include at least two consultants in the subspecialty with active and credible expertise in the relevant field of whom at least one must be a consultant paediatrician.  5. The patient is a child and I understand the Summary of Product Characteristics (SPC) states that 'there is no experience with treatment of paediatric patients below 2 years of age' and 'there is limited data in imatinib-resistant or intolerant paediatric patients below 6 years of age'.  6. Treatment with nilotinib will be as monotherapy and with dosing as described in the Summary of Product Characteristics (SPC).  7. The prescribing clinician understands the SPC cautions that in paediatric patients under nilotinib treatment is therefore recommended.  8. When a treatment break of more than 6 weeks beyond the expected cycle length is needed, I will complete a treatment break approval form to restart treatment, including indicating as appropriate if the patient had an extended break because of COVID19.  9. Nilotinib will be rewise be used as outlined in the Summary of Product Characteristics (SPC).	No	As referenced in TA425	21-Dec-16	21-Mar-17

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
NIR1	Niraparib	Niraparib as maintenance treatment in patients with high grade epithelial ovarian fallopian tube or primary peritoneal carcinoma who have a deleterious or suspected deleterious germline and/or somatic BRCA mutation and who have a recent FIRST RELAPSE of platinum-sensitive disease and who are now in response following a SECOND platinum-based chemotherapy where the following criteria have been met:  There is a separate form (NIR2) for niraparib as maintenance treatment in patients with high grade epithelial ovarian fallopian tube or primary peritoneal carcinoma who do NOT have a deleterious or suspected deleterious germline and/or somatic BRCA mutation and who are in response following platinum-based SECOND or subsequent line chemotherapy.	- achieved a complete response at the end of the 2nd platinum-based chemotherapy i.e. has no measurable or non-measurable disease on the post-chemotherapy scan and the CA125 is normal - achieved a partial response at the end of the 2nd platinum-based chemotherapy i.e. has had a at least 30% required in measurable or non-measurable disease from the start of to the completion of the 2nd platinum-based chemotherapy or the patient has a complete remission on the post-chemotherapy or T scan but the CA125 has not decreased to within the normal range.  9. The patient is currently less than 8 weeks from the date of the last infusion of the last cycle of the 2nd platinum-based chemotherapy.  10. The patient has not previously received any PARP inhibitor unless olaparib or rucaparib via the CDF has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease prompters inners in the complete of the complete of the complete or non-measurable disease prompters.	No	TA784	20-Apr-22	19-Jul-22

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
NIR2	Niraparib	Niraparib as maintenance treatment in patients with high grade epithelial ovarian, fallopian tube or primary peritoneal carcinoma who do NOT have a deleterious or suspected deleterious germline and/or somatic BRCA mutation and who have a recent FIRST OR SUBSEQUENT relapse of platinum-sensitive disease and who are now in response following a SECOND OR SUBSEQUENT platinum-based chemotherapy where the following criteria have been met:  There is a separate form (NIR1) for niraparib as maintenance treatment in patients with high grade epithelial ovarian, fallopian tube or primary peritoneal carcinoma who have a deleterious or suspected deleterious germline and/or somatic BRCA mutation and who are in response following a platinum-based SECOND line chemotherapy.	1. This application is made by and the first cycle of systemic anti-cancer therapy.  2. This patient has a proven histological diagnosis of predominantly high grade serous or high grade endometrioid or high grade clear cell ovarian, fallopian tube or primary peritoneal carcinoma.  Please enter below as to which is the predominant histology in this patient:  - high grade des records denocarcinoma or  - high grade des records denocarcinoma or  - high grade des records descreated and concerns on a concerns of the patient of the	No	TA784	20-Apr-22	19-Jul-22

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
NIV1	Nivolumab	Nivolumab for previously treated advanced renal cell carcinoma	1. This application is being made by and the first cycle of systemic anti-cancer therapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has unrescatable locally advanced or metastatic renal cell carcinoma (RCC) which has either a clear cell component or is one of the types of RCC as indicated below. Please indicate below with RCC historically applies to this patient:  8.RC with a clear cell component or Papillary RCC or  Chromophobe RCC or  Chromophobe RCC or  Chromophobe RCC or  Multilous and spindle cell RCC or  Multilous and spindle cell RCC or  Multilous and spindle cell RCC or  Multilous and spindle cell RCC or  Multilous and spindle cell RCC or  Multilous and spindle cell RCC or  Multilous and spindle cell RCC or  Multilous and spindle cell RCC or  Multilous and spindle cell RCC or  Multilous and spindle cell RCC or  Multilous and spindle cell RCC or  Multilous and spindle cell RCC or  Multilous and spindle cell RCC or  Multilous and spindle cell RCC or  Multilous and spindle RCC  3. The patient has been previously treated with only 1 or 2 previous lines of antiangiogenic therapy for advanced or metastatic disease. Please indicate below the number of prior lines of antiangiogenic therapy with which the patient has been treated:  1. prior line  2. prior lines  4. The patient is either completely treatment nalve for immune-modulatory therapies (anti-Programmed Death*-Programmed Death*-Ligand-1 (PD-1), anti-Programmed Death*-Ligand-1 (PD-1), an	No	TA417	23-Nov-16	23-Dec-16

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
NIV2	Nivolumab	The treatment of relapsed or refractory classical Hodgkin Lymphoma in ADULT patients where all the following criteria are met:	1. An application has been made by and the first cycle of systemic anti-cancer therapy is to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The prescribing clinician is aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies and hepatitis  3. The patient has a histologically confirmed diagnosis of classical Hodgkin's Lymphoma  4. The patient has released or refractory disease  5. The patient has released prior high-dose conditioning chemotherapy followed by autologous stem cell transplant (ASCT) as part of previous therapy for classical Hodgkin's Lymphoma  6. The patient has had prior treatment with brentuximab vedotin  7. The patient has nat ECOG performance status (PS) 0-1  8. The patient is an adult*  *note there is a separate Blueteq form to be used for nivolumab in this indication in children.  9. Nivolumab will be given as monotherapy.  10. The patient has no known central nervous system lymphoma.  11. The patient has not received prior treatment with an anti-PD-1, anti-PD-12, anti-CD137, or anti-Cytotoxic T-lymphocyte-associated antigen-4 (CTLA-4) antibody unless received as part of the nivolumab EAMS programme for this indication and meeting all other criteria listed.  12. The patient will receive a maximum treatment duration of 2 years of uninterrupted treatment or 52 administrations (where administered every 2 weeks) or 26 administrations (where administered every 4 weeks) with nivolumab, whichever is the later.  13. When a treatment break of more than 12 weeks beyond the expected cycle length is needed, a treatment break approval form will be completed to restart treatment.	Yes	TA462	26-Aug-17	26-Aug-1
			14. Nivolumab will otherwise be used as set out in its Summary of Product Characteristics (SPC)*  * Nivolumab can also be administered as 480mg every 4 weeks  1. An application has been made by and the first cycle of systemic anti-cancer therapy is to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The prescribing clinician is aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies and hepatitis.  3. The patient has a histologically confirmed diagnosis of classical Hodgkin's Lymphoma				
NIV3	Nivolumab	The treatment of relapsed or refractory classical Hodgkin Lymphoma in PAEDIATRIC patients where all the following criteria are met:	4. The patient has relapsed or refractory disease  5. The patient has received prior high-dose conditioning chemotherapy followed by autologous stem cell transplant (ASCT) as part of previous therapy for classical Hodgkin's Lymphoma  6. The patient has had prior treatment with brentuximab vedotin  7. The patient has an ECOG performance status (PS) 0-1  8. The patient is a child* and either post pubescent or is pre pubescent and will receive nivolumab dosage as described in the publication Blood 2016; 128: 5414  **note there is a separate Bluteq form to be used for nivolumab in this indication in adults.  9. Nivolumab will be given as monotherapy.	Yes	-	26-Aug-17	26-Aug-17
			10. The patient has no known central nervous system lymphoma.  11. Nivolumab will only be requested by and administered in principal treatment centres.  12. The use of the nivolumab has been discussed at a multi disciplinary team (MDT) meeting which must include at least two consultants in the subspecialty with active and credible expertise in the relevant field of whom at least one must be a consultant paediatrician. The MDT should include a paediatric pharmacist and other professional groups appropriate to the disease area.  13. I confirm that Trust policy regarding unlicensed treatments has been followed as nivolumab is not licensed in this indication in children.  14. The patient has not received prior treatment with an anti-PD-1, anti-PD-12, anti-CD137, or anti-Cytotoxic T-lymphocyte-associated antigen-4 (CTLA-4) antibody.  15. The patient will receive a maximum treatment duration of 2 years of uninterrupted treatment or 52 administrations (where administered every 2 weeks) or 26 administrations (where administered every 4 weeks) with nivolumab, whichever is the later.  16. When a treatment break of more than 12 weeks beyond the expected cycle length is needed, a treatment break approval form will be completed to restart treatment.				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. An application has been made by and the first cycle of systemic anti-cancer therapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
NIV4	Nivolumab	Nivolumab monotherapy for the treatment of PD-L1 positive NON-SQUAMOUS locally advanced or metastatic disease non-small cell lung cancer after chemotherapy where the following criteria have been met:	2. The patient has a histologically or cytologically confirmed diagnosis of non-squamous non-small cell lung cancer (NSCLC).  4. The patient has a histologically or cytologically confirmed diagnosis of non-squamous non-small cell lung cancer (NSCLC).  4. The patient has stage IIIB or IIIC or IV NSCLC or disease that recurred after previous potentially curative local management of NSCLC with surgery/chemoradiotherapy/radiotherapy.  5. An approved and validated test has shown that the patient's tumour expresses PD-L1 with a positive tumour proprises and an validated test has shown that the patient's tumour expresses PD-L1 with a positive tumour proprises and consider that the patient's tumour expresses PD-L1 with a positive tumour proprises and considerable provided and validated test than the patient has progressed within 6 months of completing platinum-based adjuvant or neoadjuvant therapy or chemoradiation and if appropriate that the patient has had all appropriate targeted treatments if the patient has a tumour which is positive for an actionable genomic change in relation to EGRR and KI or ROS1 or MET exon 1 aor KRAS 6312 or RET or BRAF V600 status.  7. The patient has not received prior treatment with an anti PD-1, anti-PD-12, ant		TA713	07-Jul-21	05-Oct-21
			8. Treatment with nivolumab will continue for a total of 2 years* or until disease progression or unacceptable toxicity or withdrawal of patient consent, whichever occurs first.  *2 years treatment is defined as a maximum of 52 x 2-weekly nivolumab administrations or 26 x 4-weekly administrations.  9. Nivolumab will be administered as monotherapy at a dose of 240mg every 2 weeks or 480mg every 4 weeks.  Note: nivolumab 480mg every 4 weeks is unilicensed, therefore Trust policy regarding the use of unilicensed treatments must be followed if using this dosing schedule.	late if			
			10. The patient has an ECOG performance status of 0 or 1.				
			11. The patient has no symptomatically active brain metastases or leptomeningeal metastases.				1
			12. A formal review as to whether treatment with nivolumab should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment.				
			13. When a treatment break of more than 12 weeks beyond the expected 2 or 4-weekly cycle length is needed, the prescribing clinician will complete a treatment break approval form to restart treatment, including as appropriate if the patient had an extended break on account of Covid-19.				
			14. Nivolumab will be otherwise used as set out in its Summary of Product Characteristics (SPC).				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
NIV5	Nivolumab	Nivolumab monotherapy for the treatment of SQUAMOUS locally advanced or metastatic non-small cell lung cancer after chemotherapy where the following criteria have been met:	1. An application has been made by and the first cycle of systemic anti-cancer therapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a histologically or cytologically confirmed diagnosis of squamous non-small cell lung cancer (NSCLC).  3. The patient has stage IIIB or IIIC or IV NSCLC or disease that recurred after previous potentially curative local management of NSCLC with surgery/chemoradiotherapy/radiotherapy.  4. PD-L1 testing with an approved and validated test to determine the Tumour Proportion Score (TPS) has been attempted prior to this application and the result is set out below.  Please document the actual TPS bedow (if negative, record '0) or enter 'n' ali 'the TPS cannot be documented and the reason why below:  IF A., please indicate below the reason why the actual TPS cannot be documented:	Yes	TA6SS	21-Oct-20	19-Jan-21
			7. Treatment with nivolumab will continue for a total of 2 years* or until disease progression or unacceptable toxicity or withdrawal of patient consent, whichever occurs first.  *2 years treatment is defined as a maximum of 52 x 2-weekly nivolumab administrations or 26 x 4-weekly administrations.  8. The patient will receive the licensed* dose, frequency, and route of nivolumab for this indication, as shown below  *subcutaneously—at a dose of 600mg every 2 weeks, or 1200mg every 4 weeks  *Intravenously—at a dose of 240mg every 2 weeks, or 480mg every 4 weeks (*4 weekly IV dosing is unlicensed).  9. The patient has an ECOG performance status of 0 or 1.  10. The patient has no symptomatically active brain metastases or leptomeningeal metastases.  11. When a treatment break of more than 12 weeks beyond the expected 2- or 4-weekly cycle length is needed, I will complete a treatment break approval form requesting a restart of treatment. This must be approved before nivolumab is re-commenced.				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
NIV6	Nivolumab	The treatment of recurrent or metastatic squamous-cell carcinoma of the head and neck after platinum-based chemotherapy where all the following crtieria are met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy.  2. The patient has a histologically or cytologically confirmed diagnosis of <b>squamous</b> cell carcinoma of the head and neck.  3. The patient has recurrent or metastatic head and neck cancer that is not amenable to local therapy with curative intent (surgery and/or radiation therapy with or without chemotherapy).  4. The patient's disease has progressed or recurred during or within 6 months of the last dose of previously received platinum-based chemotherapy.  Please indicate below in which disease setting this previous platinum-based chemotherapy was given:  - in the adjuvant setting or  - in the adjuvant setting or  - concurrently with radiotherapy or  - concurrently with radiotherapy or  - concurrently with radiotherapy or  - concurrently with radiotherapy or  - concurrently with radiotherapy or  - in the patient has not received prior treatment with an anti-PD-1, anti-PD-12, anti-PD-12, or anti-Cytotoxic T-lymphocyte-associated antigen-4 (CTLA-4) antibody.  7. Every effort has been made for the patient to have PD-11 testing with an approved and validated test to determine the Tumour Proportion Score (TPS). Please document the TPS results below:  1PS result on tissue (if negative enter zero):  - The TPS cannot be quantified  - PD-11 testing was not possible as the pathologist has documented that these was insufficient tissue  Please explain why TPS could not be provided:  - Subcutaneously—at a dose of 500mg every 2 weeks, or 480mg every 4 weeks (14 weekly IV dosing is unlicensed)  - Interpation tissue of 100 mg every 2 weeks, or 480mg every 4 weeks (14 weekly IV dosing is unlicensed)  - The patient will receive the licensed* dose, frequency, and route of nivolumab for this indication, as shown below  - Subcutaneously—at a dose of 500mg every 2 weeks, or 480mg every 4 weeks (14 weekly IV dosing is unlicensed)  1. The patient has no symptomatically active brain metastases or leptomeningeal metastases.  1. The patient has no	No	TA736	20-Oct-21	18-Jan-22

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
NIV7	Nivolumab	Nivolumab for the adjuvant treatment of newly diagnosed and completely resected stage IV malignant melanoma where the following criteria are met:	1. This papilication is made by and the first cycle of systemic anti-cancer therapy with nivolumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. This patient has a confirmed histological diagnosis of malignant melanoma. Please indicate whether the melanoma is BRAF V600 mutation positive or not:  8.RAF V600 mutation positive or  8.RAF V600 mutat	No	TA684	17-Mar-21	15-Jun-21

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
Blueteq Form ref:	Drug	Nivolumab monotherapy (with or without initial combination treatment with ipilimumab) for treating unresectable or advanced malignant medianoma (form a): REGISTRATION OF START OF INVOLUMAB MONOTHERAPY COMMENCED AND CURRENTLY COMMENCED AND CURRENTLY COMMENCED AND CURRENTLY CONTINUED INVOLUMAB MONOTHERAPY (WITHOUT INITIAL COMBINATION WITH IPILIMUMAB) OR OF PREVIOUSLY COMMENCED AND CURRENTLY CONTINUED NIVOLUMAB MONOTHERAPY AFTER INITIAL COMBINATION WITH IPILIMUMAB (clinicians starting patients on nivolumab plus ipilimumab combination treatment should only use this form after the ipilimumab part of the treatment has been completed). This form comes in 3 parts  1. The first part is for patients who are either scheduled to commence nivolumab monotherapy or who continue to receive nivolumab monotherapy or who commenced and continue to receive involumab monotherapy after initial combination treatment with ipilimumab. The second part of the form which must use the same unique Blueteq identifier is for those benefitting patients who choose to electively discontinue nivolumab after 2 or more years of treatment.  2. The second part (patient details will be automatically entered) will only appear once the first part of the form is approved and should be completed at the time of elective discontinuation of nivolumab. The third part of the form which must use the same unique Blueted identifier is for the same unique Blueted identifier is for the same unique Blueted identifier is for the same unique Blueted identifier is for the same unique Blueted identifier is for these	1. This application has been made by and the first cycle of systemic anti-cancer therapy with nivolumab will be/was prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy. Note: if treatment with nivolumab has already commenced, it is vital that the first treatment start date has been entered in the box above.  2. The patient has an histologically- or cyclogically-confirmed diagnosis of malignant melanoma.  3. The patient has unresectable or advanced melanoma.  4. In espect of his/her treatment for unresectable/placed disease and at the time of starting involumab, the patient his/has treatment and ipilinumah monotherapy, or staskands provisory only received BBAFMEK-largered therapy or pilinimumah monotherapy, and the standard provisory only received BBAFMEK-largered therapy or pilinimumah monotherapy, as a diagnosis of useal melanoma, and has received treatment with tebentafusp in the first line setting, and has stopped this therapy due to disease progression, or lack of tolerance  5. At the time of commencing nivolumab the patient has/had not received prior treatment with any of the following: anti-PD-1, anti-PD-12 and anti-CD137 treatments unless the patient has received adjuvant immunotherapy with nivolumab or pembrotizumab or pem	drug/	TA384 & TA400	NICE	baseline funding
		patients registered as having electively and previously stopped nivolumab and in whom there is disease progression for which the clinician wishes to recommence nivolumab monotherapy.  3. The third part of the form (patient details will be automatically entered) will only appear once the second part of the form has been approved.	intravenously – at a dose of 240mg every 2 weeks, or 480mg every 4 weeks  • 480mg IV every 8 weeks ONLY if the patient is participating in the REFINE trial (NIHR SPMS 50169).  10. When a treatment break of more than 12 weeks beyond the expected 2- or 4-weekly cycle length is needed, I will complete a treatment break approval form requesting a restart of treatment. This must be approved before will involumab is re-commenced				
			Form b and c are shown on the next page				

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
NIV8b	Nivolumab	Nivolumab for treating unresectable or advanced malignant melanoma (form b): REGISTRATION OF DISCONTINUATION OF THE OF TH	1. This registration of electively discontinued treatment with nivolumab has been made by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient is in a stable disease or a response state in relation to treatment with nivolumab for his/her melanoma.  Please indicate the nature of the response to nivolumab and if in a complete or partial response, please enter the date that this response was achieved:  - complete response and date of complete response (dd/mm/yyyy) or  - partial response and date of partial response (dd/mm/yyyy) or  - stable disease  3. The patient has either received 2 or more years of nivolumab (including any doses given with ipilimumab) or the patient was randomised to the 1 year discontinuation arm in the DANTE trial.  Please state which of these 2 reasons apply for discontinuation of therapy:  - Completed 2 or more years of nivolumab or  - Drew 1 year treatment arm in DANTE trial  Please also state the duration of treatment with nivolumab (i.e. the time between treatment commencement and discontinuation)  4. The patient has chosen this option of discontinuing therapy after an informed consenting process which has fully described the advantages and disadvantages of the options of either continuing on nivolumab or electively discontinuing nivolumab with the option of re-starting nivolumab if the disease progresses but only with nivolumab directly as the next systemic therapy following previous discontinuation of nivolumab	No	TA384 & TA400	18-Feb-16 & 27-Jul 16	18-May-16 (Blueteq approval required from 01-Feb-19)
NIV8c	Nivolumab	Nivolumab for treating unresectable or advanced malignant melanoma (form c): RE-START OF NIVOLUMAB MONOTHERAPY The third part of the form which must use the same unique Blueteq identifier is for those patients registered as having electively and previously stopped nivolumab and in whom there is disease progression for which the clinician wishes to re-commence nivolumab as the next systemic treatment.	1. This application to re-start nivolumab monotherapy has been made by and the first cycle of systemic anti-cancer therapy with nivolumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has progressive non-resectable or metastatic melanoma.  Please state the duration of time off treatment (i.e. the time between previous nivolumab discontinuation and decision to re-start nivolumab)  3. The patient has received no other systemic therapy in the time between the date of elective discontinuation of nivolumab and this application to re-start nivolumab  4. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies and hepatitis.  5. The present intention is that the patient will be treated with nivolumab monotherapy until there is progressive disease or unacceptable toxicity or if the patient declines further therapy.  6. The patient has a sufficient performance status (PS) to be fit to receive treatment with immunotherapy.  7. Nivolumab will be administered as monotherapy.  8. The licensed dose and frequency of nivolumab plus ipilimumab is not commissioned.  8. The licensed dose and frequency of nivolumab will be used (i.e. either 240mg every 2 weeks or 480mg every 4 weeks)  9. A formal medical review to assess the tolerability of treatment with nivolumab will be scheduled to occur at least by the start of the 3rd month of treatment and thereafter on a regular basis.  10. Treatment breaks of up to 12 weeks beyond the expected cycle length are allowed but solely to allow any toxicities to settle	No	TA384 & TA400	18-Feb-16 & 27-Jul 16	18-May-16 (Blueteq approval required from 01-Feb-19)

3lueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with the combination of nivolumab and ipilimumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has unresectable locally advanced or metastatic renal cell carcinoma (RCC) which has either a clear cell component or is one of the types of RCC as indicated below. Please Indicates below which RCC histology applies to this patient:  - RCC with a clear cell component or  - Papillary RCC or  - Collecting duct RCC (Bellini collecting duct RCC) or  - Medullary RCC or  - Muclinous tubular and spindle cell RCC or  - Multilocular cystic RCC or  - Which as intermediate or poor risk advanced renal cell carcinoma as assessed by the International Metastatic RCC Database Consortium (IMDC) system which scores 1 point for each of the following 6 factors – a score of 0 indicates good risk disease, a score of 1-2 indicates intermediate risk and a score of 3-6 denotes poor risk:  The IMDC factors are:  - less than 1 year from time of initial diagnosis of RCC to now  - A Karnofsky performance status of <80% (see below for description of Karnofsky scale of performance status)				
NIV9		For the 1st line treatment of intermediate or poor risk advanced renal cell carcinoma where the following criteria are met:	the haemoglobin level is less than the lower limit of normal - the corrected calcium level is >2.5mmol/L - the platelet count is greater than the upper limit of normal - the absolute neutrophil count is greater than the upper limit of normal the absolute neutrophil count is greater than the upper limit of normal Please indicate below whether the patient is in the intermediate or poor risk prognostic group Intermediate risk disease (IMDC score of 1 or 2) or - poor risk disease (IMDC score of 1 or 2) or - poor risk disease (IMDC score of 3 or 3) or 3) or 3) - poor risk disease (IMDC score of 3 or 3) or 3)	No	TA780	23-Mar-22	21-Jun-22
			5. The patient has a Karnofsky performance status of at least 70%.  The relevant part of the Karnofsky performance status scale is as follows:  100% Normal, no complaints. No signs or symptoms of disease.  90% Able to carry on normal activities. Minor signs or symptoms of disease.  80% Normal activity with effort. Some signs or symptoms of disease.  70% Cares for self. Unable to carry on normal activity or to do active work.  60% Requires occasional assistance, but is able to care for most personal needs.  50% Requires oscasional assistance and frequent medical care.  6. The patient has no symptomatic brain metastases or leptomeningeal metastases currently requiring steroids for symptom control.  7. The patient is to be treated until loss of clinical benefit or excessive toxicity or patient choice, whichever is the sooner.				
			Note: there is no stopping rule as to the maximum treatment duration of nivolumab in this indication.  8. Ipilimumab will be used at the RCC ipilimumab dose of 1mg/Kg every 3 weeks for a maximum of four 3-weekly cycles.  9. Nivolumab will be used at a dose of 3mg/Kg IV every 3 weeks for the first 4 cycles (i.e. when in combination with ipilimumab) and then as subsequent monotherapy at the licensed dose, frequency, and route for this indication, as shown below  • Subcutaneously—at a dose of 600mg every 2 weeks, or 1200mg every 4 weeks  • Intravenously—at a dose of 240mg every 2 weeks, or 480mg every 4 weeks  • 1480mg IV every 8 weeks, ONLY if the patient is participating in the REFINE trial (NIHR CPMS ID 50169).  10. Nivolumab and ipilimumab will be prescribed and administered as outlined in their respective Summary of Product Characteristics (SPCs) for this indication.  11. When a treatment break of more than 12 weeks beyond the expected 2-, 3-, or 4-weekly cycle length is needed, I will complete a treatment break approval form requesting a restart of treatment. This must be approved before ipilimumab and/or nivolumab are re-commenced  12. If the disease progresses on the nivolumab plus ipilimumab combination the next set of treatment options are those drugs which are routinely commissioned as first to be used VEGF- or VEGFR-targeting drugs ie one choice of the following: cabozantinib or pazopanib or tivozanib or sunitinib.				

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
NIV10	Nivolumab and ipilimumab	For patients with microsatellite instability high (MSI-H) or mismatch repair deficient (dMMR) metastatic or locally advanced and inoperable colorectal cancer after prior fluoropyrimidine-based chemotherapy for metastatic disease where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with nivolumab plus ipilimumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has either metastatic or locally advanced and inoperable colorectal carcinoma.  3. The patient's tumour has a documented presence of microsatellite instability-high (MSi-H) or DNA mismatch repair deficiency (dMMR) confirmed by validated testing.  4. The patient's tumour has been determined to have wild type or mutant RAS status and the result is recorded below:	No	TA716	28-Jul-21	26-0ct-21

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
NIV15	Nivolumab	For the treatment of adult patients with unresectable locally advanced or recurrent or metastatic squamous cell carcinoma of the oesophagus previously treated with a fluoropyrimidine and platrum-based combination chemotherapy where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with nivolumab monotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a histologically confirmed diagnosis of squamous cell carcinoma or adenosquamous oesophageal carcinoma.  9. Separation with the period sophing agencer the patient has: 9. Separation of the cesophageal carcinoma of the cesophagea carcinoma of the cesophageal carcinoma of the c	No	TA707	15-Jun-21	13-Sep-21

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
NIV17	<b>Nivolumab</b> as adjuvant monotherapy	For patients with completely resected oscophageal or gastro-oesophageal carcinoma who have residual pathologica disease at surgery following prior neoadjuvant chemoradiotherapy where the following criteria has been met:	This application is being made by and the first cycle of systemic anti-cancer therapy.  In patient has a histologically confirmed diagnosis of oesophageal cancer (squamous or adenocarcinoma) or adenocarcinoma of the gastro-oesophageal junction.  Please mark below which histology applies to this patient: - squamous cell cardinoma of the oesophagus - adenocarcinomo of the oesophagus - adenocarcinomo of the oesophagus - adenocarcinomo of the oesophageal junction  S. In this patient the primary treatment intent at the outset of therapy was to treat with the sequence of chemoradiotherapy followed by surgical resection.  NB The marketing authorisation of nivolumab stipulates the use of prior neoadjuvant chemoradiotherapy followed by surgery and thus NICE's considerations and recommendations are aligned to this. Patients treated with neoadjuvant chemoradiotherapy are not eligible for adjuvant nivolumab. Patients who are treated with primary chemoradiotherapy and who then progress locally and have salvage surgery are not eligible for adjuvant nivolumab. Patients who are treated with primary chemoradiotherapy and who then progress locally and have salvage surgery are not eligible for adjuvant nivolumab. Patients who are treated with primary chemoradiotherapy and who then progress locally and have salvage surgery are not eligible for adjuvant nivolumab. Patients who are treated with primary chemoradiotherapy and patient has been treated with neoadjuvant chemoradiotherapy and that the concurrent chemotherapy used with the radiotherapy was platinum-based.  Please document the number of weeks since the end of the chemoradiotherapy:  ———————————————————————————————————	No	TA746	17-Nov-21	15-Feb-22
NIV18	Nivolumab and ipilimumab	Nivolumab in combination with ipilimumab for treating advanced melanoma	Alk Nicolumab is re-commenced	No	TA400	27-Jul-16	25-Oct-16

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
NIV19	Nivolumab	Nivolumab monotherapy for adjuvant treatment after complete tumour resection in adult patients with high risk muscle invasive urothelial cancer with tumour cell PD-L1 expression of 2:1% and in whom adjuvant treatment with platinum-based chemotherapy is unsuitable where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with adjuvant nivolumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a histologically documented diagnosis of muscle invasive unrothelial cancer of the bladder, useder or renal pelvis.  Please mark below the site of origin of the unrothelial cancer: - bladder  - serial polisis  3. The patient's curribulal cancer has been documented as exhibiting PD-L1 expression on 21% of fumour cells as determined by an approved and validated PD-L1 assay.  2. The patient's curribulal cancer has been documented as exhibiting PD-L1 expression on 21% of fumour cells as determined by an approved and validated PD-L1 assay.  2. The patient was treated with necadiporant chemotherapy or not: please mark below as appropriate:  - Yes, the patient was treated with necadiporant chemotherapy or not: please mark below as appropriate: - Yes, the patient had fine treative messignant chemotherapy or not: please mark below as appropriate Yes, the patient had fine treative messignant chemotherapy. The patient had fine treative messignant chemotherapy in the patient state of the treative messignant chemotherapy. The patient had fine treative messignant chemotherapy the pathological stage for the resected tumor unsure by FD or patient and not receive messignant chemotherapy. The patient had not receive message of the patient did not receive message and the patient of the muscle invasive unrothelial cancer with all surgical margins negative for tumour i.e. a 80 resection has taken place.  - The patient has not have been any necessity and the chemotherapy in the patient cells cancer with all surgical margins negative for tumour i.e. a 80 resection has been marked by a single patient of the muscle invasive unrothelial cancer with all surgical margins negative for tumour i.e. a 80 resection has been marked by a single patient by the patient of the muscle invasiv	No	TA817	10-Aug-22	08-Nov-22

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application for nivolumab in combination with ipilimumab is being made by and the first cycle of systemic anti-cancer therapy with nivolumab in combination with ipilimumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, hepatitis and skin toxicity.  3. The patient has a histologically or cytologically confirmed diagnosis of mesothelioma.  4. The mesothelioma is of pleural or non-pleural origin.  Please indicate below the site of origin of the mesothelioma in this patient:  - the pleura or - the pericardium or - the pericardium or - the pericardium or - the tunica vaginalis in the testis  5. The histological subtype of mesothelioma as to whether the mesothelioma in this patient: - the mesothelioma is of epithelioid type or mesothelioma in this patient: - the mesothelioma is of epithelioid type or mesothelioma in this patient: - the mesothelioma is of epithelioid type or on-epithelioid type or son-epithelioid (sarcomatoid or biphasic) type or - the mesothelioma is of on-epithelioid (sarcomatoid or biphasic) type or				
NIV20	<b>Nivolumab</b> in combination with ipilimumab	For treatment of unresectable malignant mesothelioma previously untreated with systemic therapy where the following criteria have been met:	6. The patient has unresectable disease. 7. The patient has not previously received any systemic therapy for mesothelioma (neither cytotoxic chemotherapy nor immunotherapy) unless the patient was started on treatment with nivolumab and ipilumumab via the EAMS scheme and all other treatment criteria on this form are fulfilled. Please mark below which of these 2 clinical scenarios applies to this patient: - The patient has not received prior systemic treatment for mesothelioma including chemotherapy, anti-PD-1, anti-PD-1, anti-PD-1, anti-PD-1, anti-CD137, or anti-cytotoxic T-lymphocyte-associated antigen-4 (CTLA-4) antibodies Received prior treatment with nivolumab and ipilumumab via EAMS scheme and all other treatment criteria on this form are fulfilled Note: patients previously treated with cytotoxic chemotherapy for mesothelioma or with immunotherapy for mesothelioma are not eligible to receive nivolumab plus ipilimumab.	No	TA818	17-Aug-22	16-Sep-22
			8. The patient has an ECOG performance status of 0 or 1.  9. The patient has an ECOG performance status of 0 or 1.  9. The patient either has no known brain metastases or if the patient has brain metastases, the patient is symptomatically stable prior to starting nivolumab in combination with ipilimumab.  10. Nivolumab will be administered at a flat dose of 360mg every 3 weeks.  11. Nivolumab will be administered at a flat dose of 360mg every 3 weeks.  12. Ipilimumab will be administered at a dose of 1 mg/Kg every 6 weeks.  13. The patient will be administered at a dose of 1 mg/Kg every 6 weeks.  13. The patient will be treated until loss of clinical benefit or excessive toxicity or patient choice to discontinue treatment or completion of 2 years of treatment (a maximum of 35 cycles of nivolumab and a maximum of 17 cycles of				
			ipilimumab), whichever is the sooner.  Note: the registration trial for this indication (checkmate/43) had a 2 year stopping rule in the trial design and NICE's assessment of clinical and cost effectiveness was based on a treatment duration of nivolumab plus ipilimumab that reflected the 2 year stopping rule in Checkmate/43.  14. A first formal medical review as to whether treatment with nivolumab in combination with ipilimumab should continue or not will be scheduled to occur at least by the end of the first 6 weeks of treatment.  15. When a treatment break of more than 12 weeks beyond the expected 6-weekly cycle length is needed, a treatment break approval form will be completed to restart treatment, including as appropriate if the patient had an extended break on account of Covid-19.  16. The next appropriate line of therapy would be platinum-based chemotherapy in combination with pemetrexed if the patient is fit enough to receive such treatment.  17. Nivolumab and ipilimumab will be used as set out in their respective Summary of Product Characteristics (SPCs).				

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
NIV21	Nivolumab in combination with platinum and fluoropyrimidine-based chemotherapy	For previously untreated unresectable advanced or recurrent or metastatic squamous cell carcinoma of the oesophagus with a tumour cell PD-L1 expression of 1% or more and a PD-L1 combined positive score of c10 where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with nivolumab in combination with chemotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a histologically- or cytologically-confirmed diagnosis of squamous cell carcinoma of the oesophagus or adenosquamous carcinoma of the oesophagus.  Please mark below which histology applies to this patient:  3. The patient has locally advanced unresectable or recurrent or metastatic disease.  4. The patient has not received any previous systemic therapy for locally advanced unresectable or recurrent or metastatic disease.  5. An approved and validated test has demonstrated that the tumour cell PD-L1 expression is 1% or more.  Please document the actual tumour cell PD-L1 expression result below:  Tumour cell PD-L1 expression %:  6. An approved and validated test has demonstrated that the tumour has a PD-L1 expression with a combined positive score (CPS) of <10.  Please document the actual PD-L1 combined positive score (CPS) below:  PD-L1 CPS:  7. The patient has not received prior treatment with any antibody which targets PD-1 or PD-L1 or PD-L2 or CD137 or OX40 or anti-cytotoxic T-lymphocyte-associated antigen-4 (CTLA-4) unless the patient discontinued or completed checkpoint inhibitor immunotherapy as part of adjuvant therapy without disease progression and at least 6 months has elapsed between the date of the last immunotherapy treatment and the date of first diagnosis of relapse with recurrent or metastatic disease.  8. The chemotherapy used in combination with nivolumab will be both platinum and fluoropyrimidine-based.  9. Nivolumab will be administered at the licensed doses shown below  * Subcutaneously – at a dose of 600mg every 2 weeks, or 480mg every 4 weeks  * Intravenously – at a dose of 600mg every 2 weeks, or 480mg every 4 weeks	No	TA865	08-Feb-23	09-May-23
			Note: Nivolumab at a dose of 360mg, 3-weekly, when given in combination with 3-weekly based chemotherapy is permitted, but this is off-label dosing, so trust procedures for off-label prescribing must be adhered to.  10. The patient has an ECOG performance status (PS) of 0 or 1 and is fit for platinum and fluoropyrimidine-based chemotherapy in combination with nivolumab.  11. The patient has no symptomatically active brain metastases or leptomeningeal metastases.  12. Nivolumab will be stopped at whichever of the following events occurs first: disease progression or unacceptable toxicity or withdrawal of patient consent or after 2 calendar years of treatment regardless of any treatment breaks.  Note: the 2 year stopping rule for nivolumab in this indication is in the marketing authorisation and its measurement as a 2 calendar year stopping rule was part of the company submission to NICE for the clinical and cost effectiveness.  13. When a treatment break of more than 12 weeks beyond the expected 2 or 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment.  14. Nivolumab will otherwise be used as set out in its Summar of Product Characteristics (SPC).				

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
NIV22	Nivolumab in combination with platinum and fluoropyrimidine-based chemotherapy	For previously untreated advanced or metastatic HER-2 negative adenocarcinomas of the stomach, gastro- oesophageal junction or oesophagus which express Po-11 with a combined positive score of 5 or more where the following criteria have been met:	1. This againstant is being made by and the first cole of systemic active cancer therapy.  2. The patient has a bristologically or cytiologically confirmed diagnosis of HER2 negative adenocarisons of the stomach or gastro-escaphageal junction or esciphages.  2. The patient has been bristologically or cytiologically confirmed diagnosis of HER2 negative adenocarisons on the stomach.  3. HER2 negative adenocarisons and the patient is the disease application or escaphageal junction or esciphageal junction or esciphageal junction or escaphageal junction and has since had disease progression - this patient was previously treated with adjacent thereological junction or escaphageal junction or escaphageal junction and underwent surgery followed junction or escaphageal jun		TA857	11-Jan-23	11-Apr-23

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
NIV23	<b>Nivolumab</b> plus chemotherapy	For the neoadjuvant treatment of adults with previously untreated UICC/AICC 8th edition stage IIA or IIB or IIIA or N2 only IIB non-small cell lung cancer and who are candidates for potentially curative surgery where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-carecer therapy with neadjuvant involumab in combination with chemotherapy will be prescribed by a consultant specialist specifically trained and accordined in the use of systems can character charactery.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, collisis, nephritis, endocrinopathies, high papers and the bow with histologically documented diagnosis of non-small cell lung cancer (NSCLC).  Pleases make below with histology applies to this patient: - squarmous NSCLC  4. The patient either has been documented as NOT having a NSCLC which harbours an EGFR 19 or 21 mutation or an ALK gene fusion or or power with no histology and a section to not test for an EGFR 19 or 22 mutation or an ALK gene fusion or proceed with histology and be been made following discussion at the Lung Cancer MDT and consideration of the relevant patient characteristics (including age and smoking status).  Please make below which option applies to this patient: - Documented as NOT having a NSCLC which harbours an EGFR 19 or 21 mutation or an ALK gene fusion Patient has squarmous NSCLC and a decision to not test for an EGFR 19 or 21 mutation or an ALK gene fusion and proceed with nivolumab has been made following discussion at the Lung Cancer MDT.  5. The clinical TMM staging has been agreed at the appropriate Lung Cancer MDT meeting to be stage IIA or IIB or I	No	TA876	22-Mar-23	20-Jun-23

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
NIV24	<b>Nivolumab</b> with ipilimumab	Nivolumab plus ipilimumab for previously untreated patients with microsatellite instability high (MSI-H) or mismatch repair deficient (MMR) metastatic or locally advanced and inoperable colorectal cancer where the following criteria have been met:	1. This application for involumab plus ipilimumab is being made by and the first cycle of systemic anti-cancer therapy with nivolumab plus ipilimumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has either metastatic or locally advanced and inoperable colorectal carcinoma and has not received any previous systemic therapy for this indication.  Note: patients may have received neoadjuwant systemic therapy for non-metastatic disease and/or adjuvant chemotherapy after surgery.  3. The patient's tumour has a documented presence of microsatellite ratisability-high (MSH-H) or DNA mismatch repair deficiency (dMMR) confirmed by validated testing.  4. The patient's tumour has been determined to have wild type or mutant RAS status and the result is recorded below:	No	TA1065	28-May-25	27-Aug-25

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
NIVREL1	Nivolumab in combination with relatimab (Opdualag *)	As first immunotherapy for treating unresectable or metastatic melanoma in patients aged 12 years or more where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to checkpoint inhibitor treatments including pneumonitis, colitis, nephritis, endocrinopathise, hepathis mycracrifist and skin toxicities.  3. The patient has unresectable stage III or stage IV histologically confirmed melanoma.  4. The patient is aged 12 years or older.  5. The patient is not received previous treatment for this indication of unresectable or metastatic melanoma with any of the following: anti-Programmed Death receptor-1 (PD-1), anti-Programmed Death-1 ligand-1 (PD-11), anti-PO-12, or anti-cytotace? I hymphocyte associated antigen-4 (anti-CTLA-4) autibodies.  Note: treatment with involumable prevalentiam is not trunded for any patients with unresectable or metastatic melanoma who have already started treatment with pembrolizumab monotherapy or nivolumab monotherapy or absolumab roles. International horizonia in the patient is completely treatment naive for systemic therapy for melanoma or has only received specifically allowed prior systemic therapy*.  *Allowed prior therapies are:  1 pipor adjuvant therapy with adjuvant nivolumab or pembrolizumab or  2) prior immune checkpoint inhibitors for the advanced disease indication only when given as part of a clinical trial either as monotherapy or in combination with ipilimumab or  3) BRAF/MEK inhibitor targeted therapies when given for the adjuvant indication or  4) BRAF/MEK inhibitor targeted therapies when given or the adjuvant indication or  4) BRAF/MEK inhibitor targeted therapies when given or the adjuvant indication or  4) BRAF/MEK inhibitor targeted therapies when given or the adjuvant indication or  4) BRAF/MEK inhibitor targeted therapies when given or the adjuvant indication or  4) BRAF/MEK inhibitor targeted therapies when given or the adjuvant indication or  4) BRAF/MEK inhibitor targeted the	No	TA950	07-Feb-24	07-May-24

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
OBI2	Obinutuzumab	Obinutuzumab in combination with chlorambucil for untreated chronic lymphocytic leukaemia where the following criteria have been met:	1. This application is being made by and the 1st cycle of systemic anti-cancer therapy. 2. The patient has a confirmed pathological diagnosis of chronic lymphocytic leukaemia. 3. The patient has a confirmed pathological diagnosis of chronic lymphocytic leukaemia. 4. The patient has NOT been previously treated for chronic lymphocytic leukaemia and has comorbidities that make full-dose fludarabine-based therapy and bendamustine-based therapy unsuitable for them, e.g. people who have comorbidities such as impaired renal function, hypertension or diabetes 5. A maximum of 6 cycles of the combination of obinutzurumab plus chlorambucil should be used 6. The patient has a performance status (PS) of 0 - 2. 7. No planned treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve).* **Requests for continuation of treatment after unplanned treatment breaks over this duration should be made via the treatment break approval process.  8. The licensed doses and frequencies of obinutzurumab and chlorambucil will be used.	No	TA343	02-Jun-15	31-Aug-15
OBIBEN1	Obinutuzumab with bendamustine	The treatment of follicular lymphoma refractory to ritusimab where the following criteria apply:	1. An application has been made by and the first cycle of systemic anti-cancer therapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a confirmed histological diagnosis of follicular lymphoma.  3. The patient has been previously treated for follicular lymphoma with rituximab-containing chemotherapy (i.e. with induction rituximab-containing chemotherapy followed if appropriate by maintenance rituximab therapy) and that the patient has either progressed during rituximab-containing induction chemotherapy or during or within 6 months of completing maintenance rituximab monotherapy.  Please indicate below whether the patient progressed during rituximab-containing induction chemotherapy or during or within 6 months of completing maintenance rituximab monotherapy.  The patient has either failed to respond to or progressed during rituximab-containing combination induction chemotherapy or  The patient has progressed during or within 6 months of completing maintenance single agent rituximab.  If the patient progressed during or within 6 months of completing maintenance single agent rituximab, please indicate how many months since completion of previous induction rituximab-containing combination chemotherapy progression occurred:  Please also indicate below whether the patient was originally treated with 1st line obinutuzumab-containing chemotherapy or not:  The patient was previously treated with 1st line obinutuzumab-containing chemotherapy or  The patient has not previously treated with 1st line obinutuzumab-containing chemotherapy.  4. The patient has not previously received treatment with bendamustine unless completed more than 2 years previously.  5. A maximum of 6 cycles of the combination of obinutuzumab plus bendamustine should be used and followed in responding patients or in those with stable disease with maintenance single agent obinutuzumab once every 2 months for a maximum of 2 years or until disease progressio	No	TA629	13-May-20	11-Aug-20

following crtieria are met:  6. I confirm that obiniturumab is to be given in combination with induction combination chemotherapy as either:  OPTION 1 - A maximum of 6 cycles if given with bendamustine.  OPTION 2 - A maximum of 8 cycles if given with CHOP (then give obinutuzumab alone for cycles 7 and 8)  OPTION 3 - A maximum of 8 cycles if given with CVP.  7. On completion of induction chemotherapy in combination with obinutuzumab, only patients having at least a documented partial response to treatment will commence maintenance therapy with single agent* obinutuzumab once every 2 months for a maximum of 2 eyers or until disease progression (whichever occurs first)  *Note patients entered including that indicat Irail may receive lenalidomide alongside obinutuzumab maintenance therapy as per the trial protocol.  **PETRAP Hase 3 evaluation of PET-guided, Response-Adapted therapy in patients with previously untreated, high tumour burden follicular lymphoma	Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
9. A formal medical review as to whether treatment with obinutuzumab in combination with chemotherapy should continue or not will be scheduled to occur at least by the end of the third cycle of treatment  10. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment, including as appropriate if the patient had an extended break on account of Covid-19.	OB11	Obinutuzumab	follicular lymphoma where all the	2. The patient has a confirmed histological diagnosis of grade 1-3a CD20-positive follicular lymphoma  3. The patient has not previously received any of the following for treatment of lymphoma: chemotherapy alone, immunotherapy alone (rituximab, oblinutuzumab) or chemotherapy in combination with immunotherapy (rituximab, oblinutuzumab).  4. The patient has been assessed according to the Follicular Lymphoma International Prognostic Index (FLIPI) and has scored a value of at least 2. Please indicate FLIPI score:  Follicular Lymphoma International prognostic Index (FLIPI) scoring system  1. Age: If < 60 years, score 0; If 2 60 years, score 1  2. Serum LDH: If in normal range, score 0; If raised above normal range, score 1  3. Haemoglobin leve: If 2 120g/L, score 0; If \$10 to 17, score 1  3. Haemoglobin leve: If 2 120g/L, score 0; If \$10 to 17, score 1  5. Number of involved nodal areas: If < 4, score 0; If \$2, \$5, score 0. Each of the following is considered a single nodal area: left cervical, right cervical, left axiliary, right axiliary, mediastinal (includes hilar, paratracheal and retrocrural areas), insenteric (includes coeliac, splenic and portal areas), para-aortic (includes coeliac, splenic and portal areas), para-aortic (includes coemina, splenic areas), left inguinal (includes left femoral area), right inguinal (includes right femoral area), other (eg epitrochlear, popiliteal areas)  5. The patient has bulky stage II disease (>7cm) or stage III disease or stage IV disease. Patients with stage I disease or non-bulky stage II disease are not eligible for obinutuzumab  6. Lonfirm that obinutuzumab is to be given in combination with induction combination chemotherapy as either:  OPTION 2 - A maximum of 6 cycles if given with dendematusine.  OPTION 3 - A maximum of 6 cycles if given with chemotherapy in combination with obinutuzumab, only patients having at least a documented partial response to treatment will commence maintenance therapy with single agent* obinutuzumab once every 2 months for a maximum of 8	No	TA513	21-Mar-18	19-Jun-18

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3lueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
OLAP1a	Olaparib in its tablet formation	For the maintenance treatment in patients with high grade epithelial stage ill or vovarian, fallopian tube or primary peritoneal carcinoma who are in response following platinum-based FRST line chemotherapy AND who have a deleterious or suspected deleterious germline and/or somatic BRCA mutation where the following criteria have been met:  THIS FORM IS FOR INITIATION OF MAINTENANCE OLAPARIB AS A SINGLE AGENT ONLY.  THIS FORM IS FOR INITIATION OF MAINTENANCE OLAPARIB TABLETS IN THIS INDICATION. A separate CDF form OLAP1B is not for those patients with	1. This application is made by and the first cycle of systemic anti-cancer therapy with olaparib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient has a proven histological diagnosis of predominantly high grade serous or high grade endometrioid or high grade clear cell ovarian, fallopian tube or primary peritoneal carcinoma.  Please enter below as to which is the predominant histology in this patient:  - high grade serous adenocarcinoma or - high grade endometrioid adenocarcinoma or - high grade endometrioid adenocarcinoma or - high grade endometrioid adenocarcinoma or - high grade endometrioid adenocarcinoma or - high grade endometrioid adenocarcinoma or - high grade clear cell carcinoma  3. This patient has had germline and/or somatic (tumour) BRCA testing.  Please enter below the type of tissue on which BRCA mutation positive and germline BRCA mutation or - proven germline BRCA mutation or or - proven germline BRCA mutation or or - proven somatic BRCA mutation positive and germline BRCA mutation pos	Yes	TA962	28-Mar-24	26-Jun-24
			9. This patient has responded to the recently completed 1st line chemotherapy and has achieved a partial or complete response to treatment according to the definitions given below and with no evidence of progressive disease on the post-treatment scan or a rising CAL25 level. Please enter below as to which response assessment applies to this patient:  - achieved a complete response at the end of 1st line chemotherapy i.e. has no measurable or non-measurable disease from the start of to the completion of 1st line chemotherapy or the patient has a complete remission on the post-chemotherapy CT scan but the CAL25 has not decreased to within the normal range.  10. The patient has not previously received any PARP inhibitor unless 1st line maintenance niraparib monotherapy has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression. Please mark below which scenario applies to this patient:  - the patient has never previously received any received any received any received any received any received any received any received any received any received any received any received and patient in the clear absence of disease progression.  11. Olaparib will be used as monotherapy.  12. Maintenance olaparib is not being administered concurrently with maintenance bevacizumab.  Please indicate below whether bevacizumab was used in combination with the 1st line chemotherapy or  - bevacizumab 15mg/kg given in combination with platinum-based chemotherapy or  - bevacizumab 15mg/kg given in combination with themotherapy or  - no bevacizumab used in combination with themotherapy or  - no bevacizumab used in combination with themotherapy or  - no bevacizumab used in combination with themotherapy or  - no bevacizumab is marked given in combination with themotherapy or  - no bevacizumab to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment or for a total treatment duration of 2 years if the patient is i	Yes	TA962	28-Mar-24	26-Jun-24

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
OLAP1b	Olaparib in its tablet formation	positive stage III or IV ovarian, fallopian tube or primary peritoneal carcinoma who responded to platinum-based FIRST line chemotherapy AND who still have stable residual disease after 2 years of olaparib maintenance therapy and who are planned to continue with maintenance olaparib where the following criteria have been met:	1. This application is made by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient has just completed 2 years of maintenance therapy with olaparib following a response to platinum-based 1st line chemotherapy for BRCA mutation positive high grade serous or endometrioid ovarian, fallopian tube or primary peritoneal carcinoma.  3. The patient has had a scan after completing 2 years of maintenance olaparib and this scan confirms the presence of stable residual disease and serial CA125 measurements also show no evidence of disease relapse. Note: if the patient is in complete remission after 2 years of maintenance olaparib, maintenance olaparib should be discontinued as per the marketing authorisation of olaparib and the NICE guidance.  4. The prescribing clinician considers that the patient is likely to benefit from continuing on maintenance olaparib.  5. The patient continues to have a sufficiently good ECOG performance to continue on olaparib maintenance therapy.  6. Olaparib is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment.  7. Olaparib will continue to be used as monotherapy.  8. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, the prescribing clinician will complete a treatment break approval form to restart treatment.  9. Olaparib in its tablet formulation is to be otherwise used as set out in its Summary of Product Characteristics	Yes	TA962	28-Mar-24	26-Jun-24

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
OLAP2	<b>Olaparib</b> in its tablet formation	For the maintenance treatment in patients with high grade epithelial ovarian, fallopian tube or primary peritoneal carcinoma who HAVE a deleterious or suspected deleterious germline and/or somatic BRCA mutation and who have a recent FIRST RELAPSE of platinum-sensitive disease and who are now in response following a SECOND platinum-based chemotherapy where the following criteria have been met:  There is a separate form OLAP1 for olaparib in its tablet formulation as maintenance treatment in patients with high grade epithelial stage ill or V ovarian, fallopian tube or primary peritoneal carcinoma who have a deleterious or suspected deleterious memica sensitive in the substantial stage ill or V ovarian, fallopian tube or primary peritoneal carcinoma who have a five sepsion sit stage in or V ovarian, fallopian tube or primary peritoneal carcinoma who have a deleterious or suspected deleterious germline and/or somatic RBCA mutation who are in response following platinum-based THIRD or subsequent line Chemotherapy.	- achieved a partial response at the end of the 2nd platinum-based chemotherapy i.e. has had a ≥30% reduction in measurable or non-measurable disease from the start of to the completion of the 2nd platinum-based chemotherapy or the patient has a complete remission on the post-chemotherapy CT scan but the CA125 has not decreased to within the normal range.  9. The patient is currently less than 8 weeks from the date of the last infusion of the last cycle of the 2nd platinum-based chemotherapy.  10. The patient has not previously received any PARP inhibitor unless either niraparib or rucaparib via the CDF has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or olaparib tablets have been received as part of an early access scheme and the patient meets all the other criteria listed here. Please mark below which of the four scenarios applies to this patient:  - the patient has never previously received a PARP inhibitor or	No	TA908	05-Jul-23	03-Oct-23
			11. Olaparib tablets will be used as monotherapy. 12. The patient has an ECOG performance status of either 0 or 1. Please enter below as to which ECOG performance status applies to this patient:  - ECOG PS 0 or - ECOG PS 1.  Note: a patient with a performance status of 2 or more is not eligible for olaparib.				
			13. Olaparib is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment.  14. A formal medical review as to whether maintenance treatment with olaparib should continue or not will be scheduled to occur at least by the start of the third cycle of treatment.  15. No treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve).  16. Olaparib in its tablet formulation is to be otherwise used as set out in its Summary of Product Characteristics.				

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
OLAP3	<b>Olaparib</b> in its tablet formation	For maintenance treatment in patients with high grade epithelial ovarian, fallopian tube or primary peritoneal carcinoma who have a deleterious or suspected deleterious germline and/or somatic RRAC mutation and who have a recent SECOND OR SUBSEQUENT relapse of platinum-sensitive disease and who are now in response following a THIRO OR SUBSEQUENT platinum-based chemotherapy where the following criteria have been met:  This OLAP3 form should also be used for patients transitioning from olaparibic capsules to olaparib tablets in this particular indication for maintenance therapy after 3rd or subsequent platinum-based chemotherapy.  There is a separate form OLAP1 for olaparib in its tablet formulation as maintenance treatment in patients with high grade epithelial stage Ill or IV ovarian fallopian tube or primary peritoneal carcinoma who have a deleterious or suspected deleterious germline and/or somatic BRCA mutation who are in response following platinum-based FIRST line chemotherapy.  There is also a separate form OLAP2 for olaparib in its tablet formulation as maintenance treatment in patients with high grade epithelial stage Ill or IV ovarian fallopian tube or primary peritoneal carcinoma who have a deleterious gremine and/or somatic BRCA mutation who are in response following platinum-based FIRST inchemotherapy.	BRCA2 mutation or both BRCA1 and BRCA2 mutations.  6. The patient had disease which was sensitive to the penultimate line of platinum-based chemotherapy (ie the disease responded to the line of platinum-based chemotherapy preceding the most recent line of platinum-based chemotherapy).  7. The patient has recently completed a further line of platinum-based themotherapy and has received a minimum of 4 cycles of platinum-based treatment. Please enter below which line of platinum-based treatment was the most recent line of platinum-based treatment. This must be 3rd line or 4th line or greater: - 3rd line or - 4th line or greater.  8. This patient has responded to the recently completed THIRD or subsequent line platinum-based chemotherapy and has achieved a partial or complete response to treatment according to the definitions given below and there is no evidence of progressive disease on the post-treatment scan or a rising CA125 level. Please enter below as to which response assessment applies to this patient: - achieved a complete response at the end of the recent 3rd or subsequent line of platinum-based chemotherapy is has no measurable or non-measurable disease on the post-chemotherapy scan and the CA125 is normal or a raineved a partial response at the end of the recent 3rd or subsequent line of platinum-based chemotherapy is has a a 330% reduction in measurable or non-measurable disease from the start of to the completion of the 2nd platinum-based chemotherapy or the patient has a complete remission on the post-chemotherapy CT scan but the CA125 has not decreased to within the normal range.	No	TA620	15-Jan-20	14-Apr-20

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
OLAP4_v1.1	Olaparib in combination with bevacizumab	As maintenance treatment in patients with high grade egithelial stage III or IV ovarian, fallopian tube or primary peritoneal carcinoma who are in response following platinum-based FIRST line chemotherapy AND whose cancer has a positive status for homologous recombination deficiency as defined by the presence of either a deleterious or suspected deleterious BRCA 1/2 germline and/or somatic mutation or genomic instability where the following criteria have been met:  There is a separate form <b>OAP1a</b> for use of <u>olaparib monotherapy</u> as maintenance treatment in patients with high grade epithelial stage III or IV ovarian, fallopian tube or primary peritoneal carcinoma where in response following platinum-based FIRST line chemotherapy AND whose cancer has the presence of a deleterious or suspected deleterious BRCA 1/2 germline and/or somatic mutation	1. The specialists before institutement or delayer in combination with beneatzours by its less greated by a consultant specialists specialists and an extracted and according in the search strategical disposes of predominately high grade service on high grade dear real evantar, fillogian table or primary performed cardrooms.  In page 1986 of the consistence of the predomination of the		TA946	17-Jan-24	16-Apr-24

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
OLAP5	Olaparib	Olaparib monotherapy as adjuvant treatment of high-risk TRIPLE NEGATIVE early breast cancer treated with necoadjuvant or adjuvant chemotherapy and definitive local therapy in patients with a deleterious or suspected deleterious germline BRCA mutation where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with objapan's with be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient has a proven histological diagnosis of triple negative breast cancer.  3. This patient has proven histological diagnosis of triple negative breast cancer.  4. This patient has been below as to which deleterious or suspected deleterious BECA or BECA 2 mutation(s).  Please enter below as to which deleterious or suspected deleterious BECA in or BECA 2 mutation (s).  Please enter below as to which deleterious or suspected deleterious BECA in or BECA 2 mutation (s).  The patient has received completed eleterious because the second or suspected deleterious because the second or suspected deleterious or suspected deleterious as BECA a reduction.  2. The patient has received completed eleterious or suspected deleterious as BECA and BECA 2 mutation (s).  3. The patient has received completed eleterious or suspected deleterious as the second of the patient was received with a secological required contains on the second of the patient was treated with a secological required contains regimen or a beginning treatment or an adjuvant cytotoxic chemotherapy containing regimen or at least a sorted with a secological required containing regimen or at least a sorted with a secological required cytotoxic chemotherapy in the function of the patient was received with a secological required product cytotoxic chemotherapy and the second with a secological required with a received price and second or an adjuvant cytotoxic chemotherapy and the second with a second price of the patient received with a second with a second price of the patient received with a second with a second price of the patient received with a second with a second price of the patient received with a second price of the patient received with a second with a second price of the patient received with a second price of a secon	No	TA886	10-May-23	08-Aug-23

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
OLAP6	Olaparib in combination with hormone therapy	As adjuvant treatment of high-risk HORMONE RECEPOR POSITIVE HER 2 NEGATIVE early breast cancer treated with neoadjuvant or adjuvant chemotherapy and definitive local therapy in patients with a deleterious or suspected deleterious germline BRCA mutation where the following criteria have been met:		No	TA886	10-May-23	08-Aug-23

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
OLAP7	Olaparib	Olaparib monotherapy for metastatic castration-resistant prostate cancer bearing germline and/or somatic BRCA 1 or 2 mutations in patients who have progressed following previous treatment with an androgen receptor targeted agent AND HAVE ALSO BEEN TREATED WITH DOCETAXEL where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with olaparib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient either has a proven histological or cytological diagnosis of adenocarcinoma of the prostate or has presented with a clinical picture consistent with metastatic prostate cancer with both widespread bone metastases radiologically typical of prostate cancer and a serum PSA of at least \$50ng/ml.  3. This patient HAS a documented germline and/or somatic deleterious or suspected deleterious BRCA 1 or BRCA 2 mutation(s).  Please enter below as to which deleterious or suspected deleterious BRCA mutation(s) the patient has  - BRCA 1 mutation or  - BRCA 2 mutation or  - BRCA 2 mutation or  - BRCA 2 mutation or  - BRCA 2 mutation or  - BRCA 2 mutation or  - BRCA 2 mutation or  - BRCA 2 mutation or  - BRCA 2 mutation or  - BRCA 2 mutation or  - BRCA 2 mutation or  - BRCA 3 mutation or  - BRCA 2 mutation or  - BRCA 4 mutation or  - BRCA 5 mutation or  - BRCA 5 mutation or  - BRCA 6 mutation or  - BRCA 6 mutation or  - BRCA 6 mutation or  - BRCA 7 mutation or  - BRCA 9 mutation or  - BRCA 1 or BRCA 9 mutation or  - BRCA 1 or BRCA 9 mutation or  - BRCA 1 or BRCA 9 mutation or  - BRCA 1 or BRCA 9 mutation or  - BRCA 1 or BRCA 9 mutation or  - BRCA 1 or BRCA 1 or BRCA 1 or BRCA 1 or BRCA 1 or BRCA 1 or BRCA 1 or BRCA 1 or BRCA 1 or BRCA 1 or BRCA 1 or BRCA 1 or BRCA 1 or BRCA 1 or BRCA 1 or	No	TA887	10-May-23	08-Aug-23
OLAP8	Olaparib	Olaparib monotherapy for metastatic castration-resistant prostate cancer bearing germline and/or somatic BRCA 1 or 2 mutations in patients who have progressed following previous treatment with an androgen receptor targeted agent AND HAVE NOT BEEN PREVIOUSLY TREATED WITH DOCETAXEL where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with olaparib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient either has a proven histological or cytological diagnosis of adenocarcinoma of the prostate or has presented with a clinical picture consistent with metastatic prostate cancer with both widespread bone metastases radiologically typical of prostate cancer and a serum PSA of at least \$0ng/ml.  3. This patient HAS a documented germline and/or somatic deleterious or suspected deleterious BRCA 1 or BRCA 2 mutation(s).  Please enter below as to which deleterious or suspected deleterious BRCA mutation(s) the patient has  - BRCA 1 mutation or  - BRCA 2 mutation or  - both BRCA1 and BRCA2 mutations  4. This patient has hormone-relapsed (castrate-resistant) metastatic prostate cancer.  5. The patient has been previously treated with ocetaxel.	No	TA887	10-May-23	08-Aug-23

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
OLAP9	Olaparib in combination with abiraterone	The treatment of metastatic hormone- relapsed (castrate-resistant) prostate cancer in patients who are treatment naive to androgen receptor inhibitors and in whom chemotherapy is not yet clinically indicated or appropriate where the following criteria have been met:	1. This application for olaparib plus abinaterone is being made by and the first cycle of systemic anti-cancer therapy with olaparib plus abinaterone will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient either has a proven histological or cyclological diagnosis of adenocarcinoma of the prostate or has presented with a clinical picture consistent with metastatic prostate cancer with both widespread bone metastases typical of prostate cancer and a serum PSA of at least 5 foligm/ln.  3. The patient has metastatic prostate cancer.  4. The patient has not been treated with chemotherapy for the hormone-relapsed (castrate-resistant) indicated or is inappropriate (contraindicated or declined by the patient).  Note: chemotherapy given for hormone-sensitive disease earlier in the treatment pathway does not exclude patients from potential access to olaparib plus abiraterone.  6. The patient has not previously received any therapy with an androgen receptor inhibitor such as enzalutamide, abiraterone, apalutamide or darolutamide at any place in the prostate cancer treatment pathway except in progressive disease at the time such androgen receptor inhibitor therapy was discontinued.  Please mark below which scenario applies to this patient:  - the patient has not previously received any therapy with an androgen receptor inhibitor such as enzalutamide, abiraterone, apalutamide or darolutamide at any place in the prostate cancer treatment pathway OR  - the patient has not previously received any therapy with an androgen receptor inhibitor therapy was discontinued.  - the patient has not previously received any therapy with an androgen receptor inhibitor therapy was discontinued.  - The patient has not received androgen receptor inhibitor therapy was discontinued.  - The patient has not received any previously PARP inhibitor therapy was discontinued.  - The patient has not received any previously PARP inhibitor therapy was discontinued.  - T	No	TA951	07-Feb-24	07-May-24

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
OLAP10	Olaparib	Olaparib as monotherapy for treatment of adults with deleterious or suspected deleterious germline BRCAI or 2 mutations who have HER-2 negative locally advanced or metastatic breast cancer previously treated with an anthracycline and a taxane in the adjuvant/neoadjuvant/advanced disease settligs and also treated with prior endocrine-based therapy if the patient has hormone-receptor positive disease where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy. 2. This patient has a proven histological diagnosis of HR 2 registive breast cancer. 3. The patient has a proven histological diagnosis of HR 2 registive breast cancer. 4. This patient has a proven histological diagnosis of HR 2 registive breast cancer. 4. This patient has a proven histological diagnosis of HR 2 registive breast cancer. 4. This patient has a diagnosis of HR 2 registive breast cancer. 4. This patient has a diagnosis of HR 2 registive breast cancer. 4. This patient has received a strength of the diagnosis of HR 2 registive breast cancer. 5. The patient has received prior chemotherapy with an anthracycline and a taxane in any of the adjuvant or necadjuvant or advanced disease settings unless these chemotherapy agents were contraindicated Please enter below as to which of the following scenarios applies to this patient: 5. The patient has received prior chemotherapy with an anthracycline and a taxane in any of the adjuvant or necadjuvant or advanced disease settings unless these chemotherapy agents were contraindicated Please enter below as to which of the following scenarios applies to this patient: 5. The patient has received prior chemotherapy with an anthracycline and a taxane in any of the adjuvant or necadjuvant or advanced disease settings or 4-chemotherapy with an anthracycline and a taxane in any of the adjuvant or necadjuvant or advanced disease settings or 4-chemotherapy with a nathracycline and a taxane in one of these indications continued by the setting of the patient has a setting or 4-chemotherapy with a nathracycline and a taxane in one of these indications continued the adjuvant or necadjuvant or advanced disease settings and the patient has received adjuvant or his patient:  5. The patient either has triple negative disease or if the patient has hormone receptor positive disease and received appropriate endocrine-based therapy or 4-chemotherapy with both an anthracycline and a taxane in one	No	TA1040	12-Feb-25	14-Mar-25

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
OS11	Osimertinib	The the second-line treatment of locally advanced or metastatic epidermal growth factor receptor 1790M mutation-positive non small cell lung cancer in adults where all the following criteria are met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with osimertinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has histological or cytological evidence of NSCLC that carries an EGFR T790M mutation based on a validated test OB there is documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC AND there is an informative circulating free DNA test result confirming the presence of an EGFR T790M mutation.  Please mark below on which basis the diagnosis of EGFR T790M mutation positive NSCLC has been made in this patient:  Histological or cytological evidence.  Documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic disease.  4. The patient has locally advanced or metastatic disease.  4. The patient's NSCLC has been documented as exhibiting an epidermal growth factor (EGFR) mutation.  5. The patient has been documented as exhibiting unequivocal evidence of a T790M mutation.  6. There is at least evidence of radiological disease progression on 1st line EGFR-targeted tyrosine kinase (TKI) therapy and there has been no further systemic anti-cancer treatment.  Please mark below on which TKI the patient has had no prior treatment with osimertinib or osimertinib has been received as adjuvant treatment for resected stages IB to N2 only IIIB NSCLC with either an EGFR exon 19 deletion or exon 21 substitution mutation and the patient did not progress whilst still receiving adjuvant osimertinib.  7. Either the patient has had no prior treatment with osimertinib or osimertinib has been received as adjuvant treatment for resected stages IB to N2 only IIIB NSCLC with either an EGFR exon 19 deletion or exon 21 substitution mutation and the patient did not progress whilst still receiving adjuvant osimertinib.  Please mark below which scenario applies to this patien	No	TA653	14-Oct-20	12-Jan-21
OSI2	Osimertinib	For the first line treatment of locally advanced or metastatic epidermal growth factor receptor mutation-positive non-small cell lung cancer in adults where the following criteria have been met:	13. Osimerinito will be used as set out in its Summary of Product Characteristics (SPC).  1. This application is being made by and the first cycle of systemic anti-cancer therapy with osimertinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has histological or cytological evidence of NSCLC that carries a sensitising EGFR mutation based on a validated test OB there is documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC AND there is an informative circulating free DNA test result confirming the presence of a sensitising EGFR mutation.  Please mark below on which basis the diagnosis of EGFR mutation positive NSCLC has been made in this patient:  - Histological or cytological evidence.  - Documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC and there is an informative circulating free DNA test result confirming the presence of a sensitising EGFR mutation.  3. The patient has locally advanced or metastatic disease.  4. The patient's NSCLC has been documented a senhibiting an epidermal growth factor (EGFR) mutation.  5. For the locally advanced/metastic disease indication, the patient has not received any revious cytotoxic chemotherapy or immunotherapy.  6. The patient has had no prior treatment with an EGFR inhibitor unless afatinib or dacomitinib or eriotinib or gefftinib has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or osimertinib has been received as adjuvant treatment for resected stages iB to N2 only IIIB NSCLC with either an EGFR exon 19 deletion or exon 21 substitution mutation and the patient did not progress whilst if incerving adjuvant osimertinib.  Please mark below which scenario applies to this patient:  - previous treatment with a EGFR inhibitor but treatmen		TA654	14-Oct-20	12-Jan-21

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
OSI3	Osimertinib	Osimertinib for adjuvant treatment in adults after complete tumour resection in patients with UIC/ALC Sth edition stage IB or stage IIB	1. This application is being made by and the first cycle of systemic anti-cancer therapy with adjuvant osimertinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a histologically documented non-small cell lung cancer (NSCLC).  3. The patient has undergone a complete resection of the NSCL (with all surgical margins negative for tumour.  4. The pathological stage determined on this patient:	No	TA1043	26-Feb-25	27-May-25

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
OSI4	Osimertinib in combination with pemetrexed and platinum- based chemotherapy	Osimertinib in combination with pemetrexed and platinum-based chemotherapy for the first line treatment of adult patients with recurrent or locally advanced or metastatic non-small cell lung cancer exhibiting pipidermal growth factor receptor exon 19 deletions or exon 21 (1858R) substitution mutations where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy. 2. The patient has a histologically or cytologically documented non-small cell lung cancer (NSCLC) that has been shown to exhibit an epidermal growth factor (EGFR) exon 19 deletion or exon 21 (1858R) substitution mutation. 8 here is documented generate by the hug MOT that the radiological appearances are in keeping with recurrent/locally advanced/metastatic NSCLC AND there is an informative circulating free DNA test result confirming the presence of an exon 21 (1858R) substitution mutation.  Please mark below on which basis the exon 19 deletion or or exon 21 substitution mutation.  Please mark below on which basis the exon 19 deletion or exon 21 substitution mutation.  1. The patient has recurrent or locally advanced or metastatic disease.  1. A for the recurrent place all years are contained to the patient of exon 21 (1858R) substitution mutation.  2. The patient has recurrent or locally advanced or metastatic disease.  1. A for the recurrent/locally advanced or metastatic disease.  1. A for the recurrent/locally advanced or metastatic disease.  1. A for the recurrent/locally advanced or metastatic disease.  1. A for the recurrent/locally advanced or metastatic disease.  1. A for the recurrent/locally advanced or metastatic disease.  1. A for the recurrent or locally advanced or metastatic disease.  1. Peace that has no prior treatment with an EGFR exon 19 deletion or exon 21 substitution mutation and the patient did not progress whilst still receiving adjuvant colimertinib.  Please mark below which scenario applies to this patient:  1. Periodically received and patient did not progress whilst still receiving adjuvant colimertinib.  Please mark below which scenario applies to this patient:  1. O fine trimbility with a EGFR hibitor progress or metastatic disease inclination, the patient is not received any previous yerotechnotrapy regimen and then with an EGFR exon 19 deletion or exon 21 substitution mutation and the patie	No	TA1060	08-May-25	05-Aug-25

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PAL1	Palbociclib (in combination with an aromatase inhibitor)	The treatment of previously untreated, hormone receptor-positive, HER2-negative, locally advanced or metastatic breast cancer	1. This application for palbocicilib in combination with an aromatase inhibitor is made by and the first cycle of systemic anti-cancer therapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The patient has histologically or cytologically documented oestrogen receptor positive and her-2 negative breast cancer  2. The patient has had no prior treatment with a CDK 4/6 inhibitor unless either ribocicilib or abemacicilib has had to be stopped within 6 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or a CDK 4/6 inhibitor has been previously received as adjuvant therapy and treatment was completed without disease progression at least 12 months prior to the first diagnosis of recurrent or metastatic disease.  Please mark below which one of these 4 scenarios applies to this patient:  - no prior treatment with a CDK 4/6 inhibitor or  - previous treatment with the 1st line CDK4/6 inhibitor or abemacicilib but treatment has had to be stopped within 6 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of progressive disease or  - previous treatment with the 1st line CDK4/6 inhibitor or dose-limiting toxicity and in the clear absence of progressive disease or  - previous treatment with the 1st line CDK4/6 inhibitor or dose-limiting toxicity and in the clear absence of progressive disease or  - previous treatment with the 1st line CDK4/6 inhibitor or dose-limiting toxicity and in the clear absence of progressive disease or  - previous treatment with the 1st line CDK4/6 inhibitor or dose-limiting toxicity and in the clear absence of progressive disease or  - previous treatment with the 1st line CDK4/6 inhibitor or dose-limiting toxicity and in the clear absence of progressive disease or  - previous treatment with the 1st line CDK4/6 inhibitor or dose-limiting toxicity and in the clear absence of progressive disease or  - previous tre	Yes	TA495	20-Dec-17	20-Mar-18
PAL2	Palbociclib in combination with fulvestrant	For hormone receptor-positive, HER2- negative, locally advanced or metastatic breast cancer where the following criteria are met:	1. This application for palbocicilb in combination with fulvestrant is being made by and the first cycle of palbocicilb plus fulvestrant will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has histologically or cytologically documented cestrogen receptor positive and HER-2 negative breast cancer.  3. The patient has metastatic breast cancer or locally advanced breast cancer which is not amenable to curative treatment.  4. The patient has metastatic breast cancer or locally advanced breast cancer which is not amenable to curative treatment.  5. The patient has metastatic breast cancer or locally advanced breast cancer which is not amenable to curative treatment.  5. The patient has metastatic breast cancer or locally advanced breast cancer which is not amenable to curative treatment.  6. The patient has metastatic breast cancer or locally advanced breast cancer which is not amenable to curative treatment.  7. The patient has necologic performance status of 0 or 1 or 2.  8. The patient has neceived previous endocrine therapy coording to one of the three populations as set out below as these are the groups on which the NICE Technology Appraisal for palbocicib plus fulvestrant focused. Please record which population the patient falls into:  1 has progressive disease with 10 cess months of completing adjuvant endocrine therapy for early breast cancer with no subsequent endocrine therapy received following disease progression or  1 has progressive disease on 1st line endocrine therapy for advanced/metastatic breast cancer with no subsequent endocrine therapy received following disease progression or  1 has progressive disease on 1st line endocrine therapy for advanced/metastatic breast cancer with no subsequent endocrine therapy received following disease progression or  2 has progressive disease on 1st line endocrine therapy in distribution or the hash on point treatment with a CDK 4/6 inhibitor unless either abenacicible in co	Yes	TA836	26-Oct-22	24-Jan-23

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is being made by and the first cycle of panitumumab in combination with FOLFIRINOX/FOLFOXIRI chemotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient has RAS wild-type metastatic or locally advanced and inoperable colorectal cancer.  3. This patient has not received previous cytotoxic chemotherapy for metastatic disease unless there has been use of previous neoadjuvant combination cytotoxic chemotherapy for potentially resectable metastatic colorectal cancer.  Please mark below whether the patient has had neoadjuvant chemotherapy or not:  - the patient has not had previous neoadjuvant cytotoxic chemotherapy for metastatic colorectal cancer or  - the patient has been treated with previous neoadjuvant cytotoxic chemotherapy for potentially resectable metastatic colorectal cancer  4. Panitumumab in this FOLFIRINOX/ FOLFOXIRI combination is being used as either 1st line treatment for metastatic colorectal cancer or as 2nd line treatment if treated with 1st line pembrolizumab for MSI-H/dMMR disease.  Please mark below in which line of therapy the patient is having panitumumab plus FOLFIRINOX/ FOLFOXIRI chemotherapy:	-			
			- panitumumab + FOLFIRINOX/ FOLFOXIRI is being used as 1st line treatment for metastatic colorectal cancer or - panitumumab + FOLFIRINOX/ FOLFOXIRI is being used as 2nd line treatment for metastatic colorectal cancer as the patient has MSI-H/dMMR disease and has been treated with 1st line pembrolizumab or 1st line nivolumab which was previously available as an Interim COVID option  5. The patient has not received prior treatment with cetuximab or panitumumab unless this was received as part of combination neoadjuvant chemotherapy for potentially resectable metastatic disease.  Patients with potentially resectable metastatic disease who have received a neoadjuvant cetuximab/panitumumab-containing combination chemotherapy with the intention of resection if the metastases become resectable, and who do not progress while on treatment with cetuximab/panitumumab but who then become unsuitable for surgery or have unsuccessful surgery, may continue treatment with the same cetuximab/panitumumab-containing combination chemotherapy.				
PAN3	Panitumumab in combination with FOLFIRINOX or FOLFOXIRI (5-fluorouracil, irinotecan and oxaliplatin) chemotherapy	For chemotherapy-naive untreated metastatic or locally advanced and inoperable colorectal cancer where the following criteria have been met:	Patients who have successful resection(s) after neoadjuvant cetuximab/panitumumab-containing combination chemotherapy for metastatic disease and who did not progress on such chemotherapy may receive cetuximab/panitumumab with subsequent first-line combination chemotherapy if they present later with progression of metastatic disease.  Please mark below the patient's treatment status in respect of previous cetuximab/panitumumab-containing neoadjuvant chemotherapy:  - the patient has not been treated with previous chemotherapy with either cetuximab or panitumumab-containing combination chemotherapy for metastatic disease or  - the only previous cetual mab/panitumumab-containing chemotherapy was with neoadjuvant treatment for potentially resectable disease which resulted in a lack of disease progression and the patient was then unable to proceed to surgery or had unsuccessful surgery or  - the only previous cetual mab/panitumumab-containing chemotherapy was with neoadjuvant treatment for potentially resectable disease which resulted in a lack of disease progression and the patient was then able to proceed to surgery but has since relapsed	Yes	TA439	29-Mar-17	27-Jun-17
			6. The prescribing clinician is aware that if this patient has BRAF V600 mutation-positive disease, the patient will be ineligible for encorafenib plus cetuximab as a subsequent line of therapy if they receive a cetuximab/panitumumab-containing regimen now as first-line therapy.  7. The prescribing clinician is aware that from 1st December 2020 an NHS England Best Value framework is in operation for cetuximab and panitumumab in first line colorectal cancer. The choice of this panitumumab -containing regimen is therefore in line with the local application of the Best Value framework for these drugs within my organisation.  8. Panitumumab will be given in combination FOLFIRINOX/FOLFOXIRI (5-fluorouracil, irinotecan and oxaliplatin in combination) chemotherapy.	-			
			9. Panitumumab in combination with FOLFIRNOX/ FOLFOXIRI chemotherapy will be given until disease progression on this regimen and that panitumumab will be discontinued when this disease progression occurs.  If the patient experiences excessive toxicity with irinotecan and/or oxaliplatin, panitumumab can be subsequently continued in combination a fluoropyrimidine without irinotecan and/or oxaliplatin until disease progression and then will be discontinued.  Note: continued use of panitumumab beyond 1st line therapy is not commissioned once disease progression has occurred with 1st line treatment.  10. Where a treatment break of more than 6 weeks beyond the expected cycle length is needed, I will complete a treatment break form to restart treatment, including an indication as appropriate if the patient had an extended break				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with panitumumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient has RAS wild-type metastatic or locally advanced and inoperable colorectal cancer.  3. This patient has not received previous cytotoxic treatment for metastatic disease unless there has been use of previous neoadjuvant combination cytotoxic chemotherapy for potentially resectable metastatic colorectal cancer.  Please mark below whether the patient has had neoadjuvant cytotoxic chemotherapy or not:  - the patient has not had previous neoadjuvant cytotoxic chemotherapy for motatiatic colorectal cancer or  - the patient has been treated with previous neoadjuvant cytotoxic chemotherapy for potentially resectable metastatic colorectal cancer  4. Panitumumab in this irinotecan-based combination is being used as either 1st line treatment for metastatic colorectal cancer or as 2nd line treatment if treated with 1st line pembrolizumab for MSi-H/dMMR disease.  Please mark below in which line of therapy the patient is having panitumumab plus an irinotecan-based combination chemotherapy:  - panitumumab + irinotecan-based chemotherapy is being used as 1st line treatment for metastatic colorectal cancer or	-			
PAN1_v1.3	Panitumumab in combination with irinotecan-based chemotherapy	For chemotherapy-naive metastatic or locally advanced and inoperable colorectal cancer where the following criteria are met:	- pantimumab + irinotecan-based chemotherapy is being used as 2nd line treatment for metastatic colorectal cancer as the patient has MS-H/dMMR disease and has been treated with 1st line pembrolizumab or 1st line nivolumab which was previously available as an Interim COVID option  5. The patient has not received prior treatment with cetuximab or panitumumab unless this was received as part of combination neoadjuvant chemotherapy for potentially resectable metastatic disease.  Patients with potentially resectable metastatic disease who have received a neoadjuvant ectuximab/panitumumab-containing combination chemotherapy with the intention of resection if the metastases become resectable, and who do not progress while on treatment with cetuximab/panitumumab-containing combination chemotherapy.  Patients who have successful resection(s) after neoadjuvant cetuximab/panitumumab-containing combination chemotherapy for metastatic disease and who did not progress on such chemotherapy may receive cetuximab/panitumumab with subsequent first-line combination chemotherapy if they present later with progression of metastatic disease.  Please mark below the patient's treatment status in respect of previous cetuximab/panitumumab-containing combination chemotherapy:  - the patient has not been treated with previous chemotherapy with either cetuximab or panitumumab-containing combination chemotherapy for metastatic disease or  - the only previous cetuximab/panitumumab-containing chemotherapy was with neoadjuvant treatment for potentially resectable disease which resulted in a lack of disease progression and the patient was then unable to proceed to surgery or have unable to proceed to surgery or have unable to proceed to surgery or have unable to proceed to surgery or have unable to proceed to surgery or have unable to proceed to surgery or have unable to proceed to surgery or have unable to proceed to surgery or have unable to proceed to surgery or have unable to proceed to surgery or have unable to proceed to surgery or have u	Yes	TA439	29-Mar-17	27-Jun-17
			surgery or had unsuccessful surgery or - the only previous cetuvimab/panithrummab-containing chemotherapy was with neoadjuvant treatment for potentially resectable disease which resulted in a lack of disease progression and the patient was then able to proceed to surgery but has since relapsed  6. The prescribing clinician is aware that if this patient has BRAF V600 mutation-positive disease, the patient will be ineligible for encorafenib plus cetuximab as a subsequent line of therapy if they receive a cetuximab/panitumumab-containing regimen now as first-line therapy.  7. The prescribing clinician is aware that from 1st December 2020 an NHS England Best Value framework is in operation for cetuximab and panitumumab in first line colorectal cancer. The choice of this panitumumab -containing regimen is therefore in line with the local application of the Best Value framework for these drugs within my organisation.	-			
			8. Panitumumab will be given in combination with irinotecan-based combination chemotherapy. 9. Panitumumab in combination with irinotecan-based chemotherapy will be given until disease progression on this regimen and that panitumumab will be discontinued when this disease progression occurs.  If the patient experiences excessive toxicity with irinotecan, panitumumab can be subsequently continued in combination with a fluoropyrimidine alone until disease progression and then will be discontinued.  Note: continued use of panitumumab beyond 1st line therapy is not commissioned once disease progression has occurred with 1st line treatment.				
			10. Where a treatment break of more than 6 weeks beyond the expected cycle length is needed, I will complete a treatment break form to restart treatment, including an indication as appropriate if the patient had an extended break because of COVID 19  11. The use of panitumumab will be as per the Summary of Product Characteristics (SPC).	-			

3lueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PAN2_v1.2	Panitumumab in combination with oxaliplatin-based chemotherapy	For chemotherapy-naive metastatic or locally advanced and inoperable colorecta cancer where the following criteria are met:	2. This application is being made by and the first cycle of systemic anti-cancer therapy with paintiumsmab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient has not received previous creations from the patient has not received previous creations from the patient has not received previous creations from the patient has not an explainable and the patient has the had receiplivant chemotherapy or not:  - the patient has not had previous necediplowant cytotoxic chemotherapy or not:  - the patient has not had previous necediplowant cytotoxic chemotherapy for potentially resectable metastatic colorectal cancer or  - the patient has not had previous necediplowant cytotoxic chemotherapy for potentially resectable metastatic colorectal cancer or  - the patient has not his previous necediplowant cytotoxic chemotherapy for potentially resectable metastatic colorectal cancer or  - the patient has not have displayed the patient of having parliturumals by an an obligation-based chemotherapy is being used as either 1st line from the static colorectal cancer or as 2nd line treatment if treated with 1st line pembrolizumab for MSI-H/dMMR disease.  - Pantiarumable - onaligistric hased chemotherapy is being used as 1st line treatment for metastatic colorectal cancer or superior colorization of the patient has not received by patient has many parliturumable patient patients.  - Patients has not received prior treatment with cetavimab parliturumab or 1st line nivolumab which was previously available as an interior COVID option  - The patient has not received prior treatment with cetavimab parliturumab but who then become unsuitable for surgery or have unsuccessful unsupery, may continue treatment with the same cetualizable chemotherapy is described in the patient of the metastasic disease.  - Patients who have successful resection(s) after necediporant cerumably/parliturumab-containing combination chemotherapy.  - Patients who have successful	Yes	TA439	29-Mar-17	27-Jun-17
PANO1	Panobinostat	Panobinostat for treating multiple myeloma after at least 2 previous treatments	11. The use of panitumumab will be as per the Summary of Product Characteristics (SPC).  nca	No	TA380	27-Jan-16	26-Apr-16
PDL1	Pegylated Liposomal Doxorubicin	The treatment of sarcomas where all the following criteria are met:	1. An application has been made by and first cycle of systemic anti-cancer therapy to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. a) Sarcoma in patients with cardiac impairment requiring an anthracycline, 1st line indication or  b) Sarcoma in patients with cardiac impairment requiring an anthracycline, 2nd line indication  3. To be used within the treating Trust's governance framework, as Pegylated Liposomal Doxorubicin is not licensed in these indications	Yes	n/a - NHS England clinical policy	-	01-Apr-21

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			1. An application has been made by and the first cycle of systemic anti-cancer therapy is to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, hepatitis and skin toxicities.				
			3. The patient has a histologically- or cytologically-confirmed diagnosis of stage IIIB or stage IIIC or stage IV non-small cell lung cancer (squamous or non-squamous).				
			4. The patient has stage IIIB or IIIC or IV NSCLC or had disease that recurred after previous potentially curative local management of NSCLC with surgery/chemoradiotherapy/radiotherapy.				
			5. An approved and validated test has shown that the patient's tumour expresses PD-L1 with a positive tumour proportion score [TPS] of at least 1%.				
			6. The patient has progressed either after treatment with at least two cycles of platinum-based doublet chemotherapy for stage IIIB or IIIC or IV or recurrent NSCLC after previous potentially curative local management or has				
			progressed within 6 months of completing platinum-based adjuvant or neoadjuvant therapy or chemoradiation and if appropriate that the patient has had all appropriate targeted treatments if the patient has a tumour which is positive for an actionable genomic change in relation to EGFR or ALK or ROS1 or MET exon 14 or KRAS G12C or RET or BRAF V600 status.				
			7. The patient has not received prior treatment with an anti PD-1, anti-PD-L1, anti-PD-L2, anti-PD-L2, anti-CD137 or anti-cytotoxic T-lymphocyte-associated antigen-4 (CTL-4) antibody unless the patient discontinued or completed checkpoint				
			inhibitor immunotherapy as part of adjuvant/neoadjuvant/neoadjuvant/neoadjuvant/neoadjuvant/neoadjuvant/neoadjuvant/neoadjuvant/saintenance therapy without disease progression and at least 6 months elapsed between the date of the last immunotherapy treatment and the date of first diagnosis of				
			relapse with recurrent or metastatic disease.				
			Note: NHS England does not commission re-treatment with checkpoint inhibitor therapy for patients who have discontinued or completed previous checkpoint inhibitor therapy for the locally advanced/metastatic indication.				
			Please mark below if the patient received previous checkpoint inhibitor therapy and in which setting:				
		1	the patient has never received any immunotherapy for NSCLC. If so, please type 'n/a' in the 'Time gap' box below or				
		Pembrolizumab monotherapy for the	the patient has previously been treated with adjuvant immunotherapy for NSCLC and discontinued immunotherapy without disease progression and at least 6 months prior to the first diagnosis of relapse. Please document in the box				
PEMB1	Pembrolizumab	treatment of PD-L1 positive locally	below the time gap in months between completion of previous adjuvant immunotherapy and first diagnosis of disease relapse or	No	T	11-Jan-17	11-Feb-17
PEINIBI	Pembrolizumab	advanced or metastatic non-small cell lung	the patent has previously seen deated with reconstruction of the mist diagnosis of reliable. Heate and discontinued immunotive apply without discost of four first and provided the mist diagnosis of reliable. Heate and discontinued immunotive apply without discost of four first and provided the mist diagnosis of reliable.	NO	TA428	11-Jan-17	11-reb-17
		cancer after chemotherapy where the following criteria are met:	box below the time gap in months between completion of previous neoadjuvant immunotherapy and first diagnosis of disease relapse or				
		rollowing criteria are met:	the patient has previously been treated with maintenance immunotherapy post chemoradiotherapy for NSCLC and discontinued immunotherapy without disease progression and at least 6 months prior to the first diagnosis of relapse. Please document in the box below the time gap in months between completion of previous maintenance immunotherapy and first diagnosis of disease relapse				
			Time gap in months after completion of previous adjuvant or neoadjuvant or maintenance checkpoint inhibitor immunotherapy and first diagnosis of disease relapse:				
			Note: the mandatory interval between the last date of administration of any prior adjuvant/neoadjuvant/maintenance immunotherapy and the date of first relapse is at least 6 months. For patients suffering a first relapse within 6-12				
			months of previous immunotherapy, clinicians should bear in mind the long elimination half-lives of immunotherapies and make individual assessments of the overall benefit/risk ratio of re-treatment with immunotherapy.				
			8. Treatment with pembrolizumab will continue for a total of 2 years* or until disease progression or unacceptable toxicity or withdrawal of patient consent, whichever occurs first.				
			*2 years treatment is defined as a maximum of 35 x 3-weekly cycles or the equivalent numbers of cycles if 6-weekly dosing is used.				
			9. Pembrolizumab will be used as monotherapy.				
			10. The patient has an ECOG performance status of 0 or 1.	d break on			
			11. The patient has no symptomatically active brain metastases or leptomeningeal metastases.				
			12. A formal medical review as to whether treatment with pembrolizumab should continue or not will occur at least by the end of the first 6 weeks of treatment.				
			13. When a treatment break of more than 12 weeks beyond the expected cycle length is needed, I will complete a treatment break approval form to restart treatment, including as appropriate if the patient had an extended break on account of COVID 19.				
			14. Pembrolizumab will be otherwise used as set out in its Summary of Product Characteristics.				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PEMB2	Pembrolizumab	Pembrolizumab monotherapy for the first line treatment of locally advanced or metastatic non-small cell lung cancer which expresses PD-L1 with a tumour proportion score of at least 50% where all the following criteria are met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with pembrolizumab monotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a histologically- or cytologically-confirmed diagnosis of non-small cell lung cancer (squamous or non-squamous).  Peases mark below which hatchogy applies to this patient:  - squamous NSCLC  - an approved and suitabled text has demonstrated that there is PD-L1 expression of at least 50% of tumour cells (the PD-L1 tumour proportion score).  - Bear of countent the hatel PD-L1 depression below.  - PD-L1 tumour proportion score.  - Either the patient has been documented as NOT having a NSCLC which harbours an IGER 19 or 21 mutation or an ALK gene fusion and proceed with pembrolizumab has been discussed with the patient during the consenting process, i.e. the patient has a squamous cell carcinoma and a decision to not test for an IGER 19 or 21 mutation or an ALK gene fusion and proceed with pembrolizumab has been discussed with the patient during the consenting process, i.e. the patient has a squamous cell carcinoma and a decision to not test for an IGER 19 or 21 mutation or an ALK gene fusion and proceed with pembrolizumab has been discussed with the patient during the consenting process, i.e. the patient has consented to be treated with an unknown EGRF/ ALK status.  - Patient has squamous SNCLC and discission not not test for an IGER 19 or 21 mutation or an ALK gene fusion and proceed with pembrolizumab has been discussed with the patient during the consenting process.  - Either the patient has some systemic phemapy for INSCLC or the patient change of the patient has not received any previous systemic therapy for INSCL or the patient has some discussed with the patient that will be patient that the patient that will be patient that the patient that will be patient that the patient that will be patient that pembrolizumab has been discussed with the patie	No	TA531	18-Jul-18	16-Oct-18
		tow *2; pro 10. 11. 12. stai	*2 years treatment is defined as a maximum of 35 x 3-weekly cycles or the equivalent numbers of cycles if 6-weekly dosing is used or is defined by the extended dosing schedule to which the patient has been randomised as per the protocol in the NIHR-approved REFINE-Lung trial (Reference NIHR133011).  10. The patient has an ECOS performance status of 0 or 1.  11. The patient has no symptomatically active brain metastases or leptomeningeal metastases.  12. When a treatment break of more than 12 weeks beyond the expected 3- or 6- weekly cycle length is needed, I will complete a treatment break approval form which must be approved <u>BEFORE</u> treatment with pembrolizumab is restarted  13. Pembrolizumab will otherwise be used as set out in its Summary of Product Characteristics.				

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lueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with pembrolizumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			2. The prescribing clinician I am fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, hepatitis and skin toxicities.				
			3. The patient is an ADULT and has histologically documented classical Hodgkin lymphoma				
			Note: there is a separate Blueteq form to be used for pembrolizumab in this indication in children.				
			4. The patient has failed at least 2 lines of chemotherapy and also failed treatment with brentuximab vedotin.				
			5. The patient has not received stem cell transplantation of any kind.				
			6. The patient is currently ineligible for stem cell transplantation.				
		The treatment of relapsed or refractory	7. The patient is EITHER potentially a candidate for future stem cell transplantation or not. Please mark appropriately in one of the boxes below:				
		classical Hodgkin lymphoma in ADULTS	- The patient is a candidate for future stem cell transplantation if there is sufficient benefit of treatment with pembrolizumab or				
PEMB5	Pembrolizumab	who are stem cell transplant-ineligible and	- The patient is not a candidate for stem cell transplantation however good the response to pembrolizumab may be	Yes	TA967	01-May-24	30-Jul-24
		have failed brentuximab vedotin where	8. The patient has an ECOG performance status (PS) of 0 or 1.				
		the following criteria have been met:	9. The patient has not received prior treatment with an anti-PD-1, anti-PD-12, anti-PD-12, anti-CD137, or anti-cytotoxic T-lymphocyte-associated antigen-4 (CTLA-4) antibody.				
			10. Pembrolizumab is being given as monotherapy and will commence at a fixed dose of either 3-weekly cycles of pembrolizumab monotherapy 200mg or 6-weekly cycles of pembrolizumab monotherapy 400mg.				
			11. A formal medical review as to whether treatment with pembrolizumab should continue or not will be scheduled to occur at least by the end of the third cycle of treatment if 3-weekly administration of pembrolizumab or by the end of the second cycle if 6-weekly administration is used.				
			12. The patient will be treated until stem cell transplantation occurs or loss of clinical benefit or excessive toxicity or patient choice to discontinue treatment or completion of 2 years of treatment with pembrolizumab, whichever is the sooner.	2			
			13. The patient will receive a maximum treatment duration with pembrolizumab of 2 years (or 35 x 3-weekly cycles of pembrolizumab or its equivalent if 6-weekly pembrolizumab dosing is used).				
			14. When a treatment break of more than 12 weeks beyond the expected 3- or 6-weekly cycle length is needed, a treatment break approval form to re-start treatment will be completed.				
			15. Pembrolizumab will otherwise be used as set out in its Summary of Product Characteristics (SPC).				
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with pembrolizumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				1
			2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, hepatitis and skin toxicities.				
			3. The patient is a CHILD aged 3 years and older and has histologically documented classical Hodgkin lymphoma.				
			Note: there is a separate Blueteq form to be used for pembrolizumab in this indication in adults.				
			4. The patient has failed at least 2 lines of chemotherapy and also failed treatment with brentuximab vedotin.				
			5. The patient has not received stem cell transplantation of any kind.				
			6. The patient is currently ineligible for stem cell transplantation.	-			
			7. The patient is EITHER potentially a candidate for future stem cell transplantation or not. Please mark appropriately in one of the boxes below:	- 1			
		The treatment of relapsed or refractory classical Hodgkin lymphoma in CHILDREN	- The patient is a candidate for future stem cell transplantation if there is sufficient benefit of treatment with pembrolizumab or				
PEMB6	Pembrolizumab	who are stem cell transplant-ineligible and	- The patient is not a candidate for stem cell transplantation however good the response to pembrolizumab may be	Yes	TA967	01-May-24	30-Jul-24
T EIVIDO	rembiolizamab	have failed brentuximab vedotin where	8. The patient has an ECOG performance status (PS) of 0 or 1 or its equivalent Lansky score.	163	1207	01-Way-24	30-341-24
		the following criteria have been met	9. The patient has not received prior treatment with an anti-PD-1, anti-PD-11, anti-PD-12, anti-CD137, or anti-cytotoxic T-lymphocyte-associated antigen-4 (CTLA-4) antibody.	1			
			10. Pembrolizumab is being given as monotherapy and will commence at a dose of 2mg/kg bodyweight up to a maximum of 200mg in 3-weekly cycles of pembrolizumab monotherapy.	-			
			11. A formal medical review as to whether treatment with pembrolizumab should continue or not will be scheduled to occur at least by the end of the third cycle of treatment with 3-weekly administration of pembrolizumab.				
			12. The patient will be treated until stem cell transplantation occurs or loss of clinical benefit or excessive toxicity or patient choice to discontinue treatment or completion of 2 years of treatment with pembrolizumab, whichever is the	2			
			sooner.  13. The patient will receive a maximum treatment duration with pembrolizumab of 2 years (or 35 x 3-weekly cycles of pembrolizumab).	† J			
			14. When a treatment break of more than 12 weeks beyond the expected 3-weekly cycle length is needed, a treatment break approval form to re-start treatment will be completed.	- I			
			1 1 1 1	4 J			
			15. Pembrolizumab will otherwise be used as set out in its Summary of Product Characteristics (SPC).			1	

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
РЕМВ7	Pembrolizumab	Pembrolizumab for adjuvant treatment of melanoma with high risk of recurrence where the following criteria have been met:	1. This application is made by and the first cycle of systemic anti-cancer therapy with pembrolizumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, hepatitis and skin toxicities.  3. This patient has a confirmed histological diagnosis of malignant melanoma Please includate whether the melanoma is BBAF VEOD mutation positive or not:  8. BRAF VEOD mutation negative  4. The patient has melanoma which has been staged as stage III disease according to the ALCC 8th edition. Please state which stage disease the patient has:  Stage III disease or or stage III disease and this has been done with either a sentinel lymph node biopsy ('sentinel lymphadenectomy') or when indicated with a completion lymph node dissection.  6. The patient is treatment naive to any systemic therapy for malignant melanoma and in particular has not previously received any immunotherapy with any check point inhibitors or BRAF V600 inhibitors or MEX inhibitors.  Note: NHS England does not commission any adjuvant immunotherapy with checkpoint inhibitors for stage III disease in patients who have previously received adjuvant immunotherapy for stage III disease.  7. The prescribing clinician has discussed with the patient the benefits and toxicities of adjuvant pembrolizumab in	No	TA766	02-Feb-22	03-May-22

ueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PEMB8	Pembrolizumab	Pembrolizumab in combination with pemetrexed- and platinum-based chemotherapy for the first line treatment of PD-L1 positive or negative locally advanced or metastatic non-squamous non-small cell lung cancer where all the following criteria are met:	1. This application has been made by and the first cycle of systemic anti-cancer therapy will be prescribed by a consultant specialist specifically trained and accerdited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonits, colitis, nephritis, endocrinospathies, personal control of the patient has a histologically- or cytologically-control diagnosis of non-squamous non-small cell lung cancer (NCCLC).  3. The patient has a histologically- or cytologically-control diagnosis of non-squamous non-small cell lung cancer (NCCLC).  5. EGFR and ALK mutation testing have been done and both are negative.  6. PD-L1 testing make gell low of IIC or VINSCLC or hist dissense that has researced faller potentially curative treatment with local management of NSCLC with surgery/chemoradiotherapy/fadiotherapy.  5. EGFR and ALK mutation testing have been done and both are negative.  6. PD-L1 testing with an approved and voil disease that has researced faller potentially curative treatment with local management of the squamous non-related and the result of the patient of the	No	TA683	10-Mar-21	08-Jun-2:
			- cisplatin OR - carboplatin (AUC 5)  10. On completion of 4 cycles of pembrolizumab plus pemetrexed with carboplatin or cisplatin based chemotherapy, pembrolizumab will be administered as 3-weekly or 6-weekly cycles or pembrolizumab will be administered according to the extended dosing schedule to which the patient has been randomised as per the protocol in the NIHR-approved REFINE-Lung trial (Reference NIHR133011).  11. On completion of 4 cycles of pembrolizumab plus pemetrexed-based chemotherapy in combination with cisplatin or carboplatin and in the absence of disease progression, treatment with pembrolizumab will continue for a total treatment duration of 2 years* or until disease progression or unacceptable toxicity or withdrawal of patient consent, whichever occurs first.  *2* years treatment is defined as a maximum of 3 x 3-weekly yoldes or the equivalent numbers of cycles if either 6-yoldes in defined by the extended dosing schedule to which the patient has been randomised as per	_			
			the protocol in the NIHR-approved REFINE-Lung trial (Reference NIHR133011).  12. The patient has a performance status (PS) of 0 or 1 and is fit for pemetrexed- and platinum-based chemotherapy in combination with pembrolizumab.  13. The patient has no symptomatically active brain metastases or leptomeningeal metastases.  14. A formal medical review as to whether treatment with pembrolizumab in combination with pemetrexed plus cisplatin/carboplatin should continue or not will be scheduled to occur at least by the end of the first 6 weeks of treatment.	-			
			15. Where a treatment break of more than 12 weeks beyond the expected cycle length is needed, a treatment break form will be completed to restart treatment, including an indication as appropriate if the patient had an extended break because of COVID 19.  16. Pembrolizumab will be otherwise used as set out in its Summary of Product Characteristics.				

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
РЕМВ9а	Pembrolizumab	Pembrolizumab monotherapy for treating unresctable or advanced malignant melanoma (forma). REGISTRATION OF START OF PEMBROLIZUMAB MONOTHERAPY OR OF PERVIOUS Y COMMENCE AND CURRENTLY CONTINUED PEMBROLIZUMAB MONOTHERAPY.  This form comes in 3 parts.  1. The first part is for patients who are either scheduled to commence pembrolizumab monotherapy or who commenced and continue to receive pembrolizumab menotherapy or who commenced and continue to receive pembrolizumab and continue to receive pembrolizumab and part of the form which must use the same unique Blueteq identifier is for those benefitting patients who choose to electively discontinua pembrolizumab after 2 or more years of treatment; this second part (patient details will be automatically entered) will only appear once the first part of the form is approved and should be completed at the time of elective discontinuation of pembrolizumab.  3. The third part of the form which must use the same unique Blueteq identifier is for those patients; registered as having electively and previously stopped pembrolizumab and in whom there is disease progression for which the clinician wishes to re-commence pembrolizumab in third part of the form is approved in which the automatically entered) will only appear once the second part of the form may be approved.		No	ТАЗ66	25-Nov-15	23-Feb-2016 (Blueteq approval required from 01-Feb-19)
РЕМВ9Ь	Pembrolizumab	Pembrolitumab monotherapy for treating unresectable or advanced malignant melanoma (form b): REGISTRATION OF DISCONTINUATION OF PEMBROLIZUMAB  This second part of the form which must use the same unique Blueteq identifier is for those patients in stable or response remission who have those to elettively discontinue pembrolizumab. The third part of the form which usus use that same unique Blueteq identifier is for those patients registered as having electrively and previously stopped pembrolizumab and in whom there is disease progression for which the clinician whiste for ecommence pembrolizumab, this third part of the form lastient details will be automatically entered will only appear once the second part of the form has been approved.	1. This registration of electively discontinued treatment with pembrolizumab has been made by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient is in a stable disease or a response state in relation to treatment with pembrolizumab for his/her melanoma.  Please indicate the nature of the response to pembrolizumab and if in a complete or partial response, please enter the date that this response was achieved:  - complete response and date of complete response (dd/mm/yyyy) or  - partial response and date of partial response (dd/mm/yyyy) or  - stable disease  3. The patient has either received 2 or more years of pembrolizumab (including any doses given with ipilimumab) or the patient was randomised to the 1-year discontinuation arm in the DANTE trial.  Please state which of these 2 reasons apply for discontinuation of therapy:  - Completed 2 or more years of pembrolizumab or  - Drew 1-year treatment arm in DANTE trial  Please also state the duration of treatment with pembrolizumab (i.e. the time between treatment commencement and discontinuation)  4. The patient has chosen this option of discontinuing therapy after an informed consenting process which has fully described the advantages and disadvantages of the options of either continuing on pembrolizumab or electively discontinuing pembrolizumab with the option of re-starting pembrolizumab if the disease progresses but only with pembrolizumab directly as the next systemic therapy following previous discontinuation of pembrolizumab  Form C is shown on the next page	No	TA366	25-Nov-15	23-Feb-2016 (Blueteq approval required from 01-Feb-19)

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PEMB9c		Pembrolizumab monotherapy for treating unresectable or advanced malignant melanoma (form c): RE-START OF PEMBROLIZUMAB MONOTHERAPY  The third part of the form which must use the same unique Blueteq identifier is for those patients registered as having electively and previously stopped pembrolizumab and in whom there is disease progression for which the clinician wishes to re-commence pembrolizumab as the next systemic treatment.	1. This application to re-start pembrolizumab has been made by and the first cycle of systemic anti-cancer therapy with pembrolizumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has progressive non-resectable or metastatic melanoma.  Please state the duration of time off treatment (i.e. the time between previous pembrolizumab discontinuation and decision to re-start pembrolizumab)  3. The patient has received no other systemic therapy in the time between the date of elective discontinuation of pembrolizumab and this application to re-start pembrolizumab  4. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies and hepatitis.  5. The present intention is that the patient will be treated with pembrolizumab until there is progressive disease or unacceptable toxicity or if the patient declines further therapy.  5. The present intention is that the patient will be treated with pembrolizumab until there is progressive disease or unacceptable toxicity or if the patient declines further therapy.  7. Pembrolizumab will be administered as monotherapy  8. The licensed dose and frequency of pembrolizumab will be used. *Can use either 3-weekly cycles of pembrolizumab monotherapy 200mg (or if the patient is stable and well, 6-weekly cycles of pembrolizumab monotherapy 400mg)  9. A formal medical review to assess the tolerability of treatment with pembrolizumab will be scheduled to occur by the start of the 3rd 3-weekly cycle of treatment (or equivalent if having 6 weekly dosing) and thereafter on a regular	No	TA366	25-Nov-15	23-Feb-2016 (Blueteq approval required from 01-Feb-19)
		ba	basis 10. Treatment breaks of up to 12 weeks beyond the expected cycle length are allowed but solely to allow any toxicities to settle				

1. This application is being made by and the first cycle of systemic anti-cancer therapy.  2. The prescribed gindical in fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, heaptitis and skin toxicities.  3. The patient has a spistologically- or cytologically-confirmed diagnosis of squamous non-small cell lung cancer (NSCLC).  4. The patient has stage lills or lill or if V NSCLC or has disease that has recurred after previous potentially curative local management of NSCLC with surgery/chemoradiotherapy/radiotherapy.  5. PD-L1 testing with an approved and validated test to determine the Turnour Proportion Score (TPS) has been attempted prior to this application and the result is set out below.  Note: for fully informed consent of all the potential 1st line treatment options, PD-L1 testing must still be attempted and recorded here.  Please document the actual TPS below (if negative, record '0') or enter 'n/a' if the TPS cannot be documented and the reason why:  TPS			nce	unding started
Please mark below which of these two scenarios applies to this patient: the PD-L1 TPS could not be documented (see criterion 5) or PD-L1 TPS of 0-49% or P	EMB10_v1.2	No TA770 09-Fe	nce s	_

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PEMB12	Pembrolizumab	For previously untreated metastatic or unresectable recurrent PP-L1 positive head and neck squamous cell carcinoma (HNSCC) where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with pembrolizumab monotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies and hepatitis.  3. The patient has a documented histological diagnosis of squamous cell carcinoma of the head and neck.  4. The patient has either metastatic head and neck cancer or locally advanced/unresectable recurrent head and neck cancer that is not amenable to curative intent with local therapy (surgery and/or radiation therapy with or without chemotherapy).  5. PD-L1 testing with an approved and validated test to determine the Combined Positive Score (CPS) has been done prior to this application and the CPS is ≥1% and the result is set out below. Please document the actual CPS below  Note: pembrolizumab is not funded in this indication for patients with tumours without a documented ≥1% positive PD-L1 CPS score.  6. The patient has not received prior treatment with an anti-PD-1, anti-PD-L2, anti-CP137, or anti-Cytotoxic T-lymphocyte-associated antigen-4 (CTLA-4) antibody unless the patient has received pembrolizumab monotherapy for this indication via Interim COVID19 funding.  Please tick one of the following options which applies as to any previous systemic therapy:  - the patient has not received pembrolizumab monotherapy as 1st line therapy for this metastatic/locally advanced/unresectable recurrent indication as part of Interim COVID19 funding  8. Pembrolizumab will only be administered as monotherapy as 1st line therapy for this metastatic/locally advanced/unresectable recurrent indication as part of Interim COVID19 funding  8. Pembrolizumab will only be administered as monotherapy as 1st line therapy for this metasta	No	TA661	25-Nov-20	23-Feb-21
PEMB14_v1.2	Pembrolizumab	either metastatic or locally advanced and inoperable colorectal cancer exhibiting	1. This application is being made by and the first cycle of systemic anti-cancer therapy with pembrolizumab monotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, hepatitis and skin toxicity.  3. The patient has either metastatic or locally advanced and inoperable colorectal carcinoma.  4. The patient's tumour has a documented presence of microsatellite instability-high (MSI-H) or DNA mismatch repair deficiency (dMMR) confirmed by validated testing.  5. Wild type or mutant RAS status has been determined on this patient's tumour and the result is recorded below:  - wild type RAS status  - mutant RAS status  - Test result not yet reported and the decision to proceed without knowing RAS status has been discussed with the patient during consenting process.  6. Wild type or mutant BRAF status has been determined on this patient's tumour and the result is recorded below:  - wild type RAS status  - Test result not yet reported and the decision to proceed without knowing BRAF status has been discussed with the patient during consenting process.  7. The patient has not received previous systemic therapy for metastatic colorectal cancer unless this was given with neoadjuvant intent.  Please mark below which clinical scenario applies to this patient:  - no previous systemic therapy for metastatic colorectal cancer and no previous neoadjuvant chemotherapy for metastatic colorectal cancer and no previous neoadjuvant chemotherapy for metastatic colorectal cancer and no previous neoadjuvant chemotherapy for metastatic colorectal cancer and no previous neoadjuvant chemotherapy for metastatic colorectal cancer and no previous neoadjuvant chemotherapy for metastatic colorectal cancer and no previous neoadj	No	TA709	23-Jun-21	21-Sep-21

lueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PEMB1S	Pembrolizumab in combination with platinum and fluoropyrimidine-based chemotherapy	For previously untreated advanced ossophageal carcinoma which expresses	Blueteq Approval Criteria  1. This application is being made by and the first cycle of systemic and cancer therapy will be personable by an consultant specialist specifically trained and accredited in the use of systemic and cancer therapy.  2. The personable of continuous following and cancer therapy will be personable to embassion with chemotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic and cancer therapy.  2. The personable of continuous fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-1 or anti-PD-1 treatments including personability. Cells, responsible, specifically trained and accreditation of the conspiculation of the cons	drug/ indication	TA737	NICE	baseline funding
			Note: once pembrolizumab is stopped after 2 years of treatment, it cannot be re-started.  13. A formal medical review as to how pembrolizumab plus chemotherapy is being tolerated and whether pembrolizumab should continue or not will be scheduled to occur at least by the end of the second 3-weekly cycle of treatment.  14. When a treatment break of more than 3 months beyond the expected 3- or 6-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment.  15. Pembrolizumab will otherwise be used as set out in its Summary of Product Characteristics (SPC) with the exception of criterion 12.				

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PEMB16	Pembrolizumab	For relapsed/refractory classical Hodgkin lymphoma in patients aged 3 years and older who have been treated with stem cell transplantation but never previously received brentumab vedotin where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-1 or anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, hepatitis and skin toxicity.  3. The patient has a histologically confirmed diagnosis of classical Hodgkin lymphoma.  4. The patient has relapsed or refractory Hodgkin lymphoma following stem cell transplantation.  Please mark below whether the patient had autologous and/or allogeneic stem cell transplantation:  - autologous transplantation only - allogeneic transplantation only - allogeneic transplantation only - both autologous and allogeneic resolution of the patient has never previously been treated with brentuvimab vedotin.  7. The patient has not received prior treatment with any antibody which targets PD-1 or PD-L1 or PD-L2 or CD137 or OX40 or anti-cytotoxic T-lymphocyte-associated antigen-4 (CTLA-4).  8. The patient has not received prior treatment with any antibody which targets PD-1 or PD-L1 or PD-L2 or CD137 or OX40 or anti-cytotoxic T-lymphocyte-associated antigen-4 (CTLA-4).  8. The patient has an ECOG performance status (PS) of O or 1 and is fit for treatment with pembrolizumab.  9. Pembrolizumab will be administered as monotherapy:  4. For padelatric patients (aged 18 years and older), at a dose of either 200mg 3-weekly or 400mg 6-weekly.  4. For padelatric patients (aged between 3 and 17 years), pembrolizumab will commence at a dose of 2mg/kg bodyweight up to a maximum of 200mg 3-weekly.  4. For padelatric patients (aged between 3 and 17 years), pembrolizumab will commence at a dose of 2mg/kg bodyweight up to a maximum of 200mg 3-weekly.  4. For padelatric patients (aged between 3 and 17 years), pembrolizumab will be scheduled to occur at least by the end of the second 3-weekly cycles or its equivalent if 6-weekly dosing is useed).  4. Note: the 2 year	No	TA772	23-Feb-22	24-May-22

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PEMB17	Pembrolizumab	Pembrolizumab monotherapy for relapsed/refractory classical Hodgkin lymphoma in patients aged 3 years and older who have NOT been previously treated with stem cell transplantation or brentuximab vedotin	1. This application is being made by and the first cycle of systemic anti-cancer therapy with pembrolizumab monotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-1 or anti-PD-L1 treatments including pneumonitis, collisis, nephritise, nedocrinopathies, hepatias and skin toxicity.  3. The patient is aged 3 years and older.  Please mark below whether the patient is aged 3-17 years or 18 years and older:  - the patient is aged between 3 and 17 years or  - the patient is aged alt years and older.  5. The patient has ged between 3 and 17 years or  - the patient is aged alt years and older.  5. The patient has relapsed or refractory Hodgkin lymphoma following 2 prior lines of cytotoxic chemotherapy.  6. The patient has never previously been treated with brentusinab vectorin.  7. The patient has not been previously treated with stem cell transplantation of any kind.  8. The patient has not been previously treated with stem cell transplantation of any kind.  8. The patient is currently ineligible for stem cell transplantation of any kind.  8. The patient is currently ineligible for stem cell transplantation if there is sufficient benefit of treatment with pembrolizumab OR is not a candidate for stem cell transplantation however good the response to pembrolizumab may be.  Please mark below the patient status as regards future autologous/allogeneic stem cell transplantation if there is sufficient benefit of treatment with pembrolizumab will be administered as monotherapy:  - the patient is a candidate for future stem cell transplantation in there is sufficient benefit of treatment with pembrolizumab  - the patient is an accordate for future stem cell transplantation in there is sufficient benefit of treatment with pembrolizumab  - the patient is an accordate for future st	No	TA772	23-Feb-22	24-May-22

Blueteq Form rel	: Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PEMB18_v1.2	Pembrolizumab in combination with paclitaxel or nab-paclitaxel	The treatment of previously untreated locally advanced unresectable or metastatic triple negative breast cancer in patients with PD-L1 expression test results of immune cell (IC) <13° and a combined positive score (CPS) of 10 or more where the following criteria have been met:	1. An application has been made by and the first cycle of systemic anti-cancer therapy with perhodicisans in combination with pacilizated or nab-pacitizated will be prescribed by a consultant specialist specifically trained and according in the use of systemic and-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocringosthies, persists and aktin tockly.  3. The patient has a histologically- or cytologically-confirmed diagnosis of breast cancer.  4. The patient has a histologically- or cytologically-confirmed diagnosis of breast cancer.  5. The patient has a histologically- or cytologically-confirmed diagnosis of breast cancer.  6. The patient is breast cancer has had receptor analysis performed and this in negative for all of the following: the HER2 receptor, oestrogen receptor and progesterone receptor i.e. the patient has triple negative disease.  6. The patient's tumour has been tested by an approved and wildidated test for PD-L1 expression as measured by the immune cell (IC) tests of the PD-L1 substance in the patient tumour has been tested by an approved and wildidated test for PD-L1 expression as measured by the combined positive score (IC) test and the result is 10 or more.  9. The patient's tumour has been tested by an approved and wildidated test for PD-L1 expression as measured by the combined positive score (IC) test and the result is 10 or more.  9. L1 expression with the ICP Stest:  1. Possible of the PD-L1 expression are required as the manufacturer of pembrolizumab, MSO, only sought a recommendation from NICE for patients who were ineligible for aterolizumab and had a PD-L1 expression test result as measured by the combined positive score (IC) test of 10 or more.  9. Either the patient has never had any prior treatment with anti-PD-L1PD-L1 therapy for the best cancer or the only previous anti-PD-L1PD-L	No	TA801	29-Jun-22	27-Sep-22

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PEM819_v1.1	Pembrolizumab	Pembrolizumab monotherapy for adjuvant treatment after complete tumour resection of renal cell carcinoma in adult patients at increased risk of recurrence following nephrectomy or following nephrectomy and resection of all metastatic disease where the following criteria have been met:	1. An application has been made by and the first cycle of systemic anti-cancer therapy with adjavant percentage (initicals is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PO-11 treatments including pneumonitis, collisis, nephritis, endocrinogatine, heapitists and skin toxicities.  3. The pattern has a histologically documented diagnosis of renal cell carcinoma (RCC).  Places indicate below with RCC histology applies to this patient:  **RCC with a celer cell component or Controllable RCC or Collecting due 18 CC or American studies and spindle cell RCC or Collecting due 18 CC or Coll	No	TA830	19-Oct-22	17-Jan-23

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PEMB20_v1.0	Pembrolizumab	Pembrolizumab for the adjuvant treatment of newly diagnosed and completely rescreted stage lilo stratege liC malignant melanoma where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with adjuvant pembrolizumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, hepatitis and skin toxicities.  3. This patient has a documented histological diagnosis of malignant melanoma.  Please indicate whether the melanoma is BRAF V600 mutation positive or not:  -BRAFV600 mutation positive or  -BRAFV600 mutation in regulative  4. The patient has melanoma which has been staged as stage IIB or stage IIC disease according to the AICC 8th edition.  Please state which stage disease the patient has:  -Stage III disease or  -Stage III disease or  -Stage III disease or  -Stage III disease or  -Stage III disease or  -Stage III disease or  -Stage III disease or  -Stage III disease or stage III disease.  6. Complete resection has taken place for stage II disease.  6. The patient is treatment naïve to any systemic therapy for malignant melanoma and in particular has not previously received any immunotherapy with any check point inhibitors or BRAF V600 inhibitors.  Note: NHS England does not commission any adjuvant immunotherapy with checkpoint inhibitors for stage III disease in patients who have previously received adjuvant immunotherapy for stage IIB or IIC disease.  7. The prescribing clinician has discussed with the patient the benefits and toxicities of adjuvant pembrolizumab in stage IIB/IIC disease and has used the expected median figures below for melanoma-specific survival in relation to the risk of disease relapse II a routine surveillance policy is followed:  -In stage III disease, the 5 and 10 year figures for melanoma-specific survival probabilities with routine surveillance are 87% and 82%, respectivel	No	TA837	26-Oct-22	24-Jan-23

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advanced or early staget triple negative breast cancer at high risk of recurrence where the following criteria have been met:  1. During the neoadjuvant phases of treatment the patient will be treated with a fixed dose of pembrolizumab or 6 cycles of 1s. 3-weekly epubrolizumab or 6 cycles of 1s. 4-weekly epubrolizumab or 6	Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
18. A formal medical review as to how pembrolizumab and neoadjuvant chemotherapy are being tolerated and whether neoadjuvant chemotherapy should continue or not will be scheduled to occur at least by the end of the first 6 weeks of treatment.  19. When a treatment break of more than 12 weeks beyond the expected 3 or 6 weekly cycle length is needed, a treatment break approval form will be completed to restart treatment.	PEMB21	Pembrolizumab	chemotherapy as neoadjuvant treatment and then continued as adjuvant monotherapy after definitive surgery for patients with previously untreated locally advanced or early stage triple negative breast cancer at high risk of recurrence where the following criteria have been	and accretited in the use of systemic anti-cancer through.  The prescripting claims is fully wave for the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PP-L1 treatments including pneumonitis, collis, nephritis, endocrinopathles, hepatitis and skin toxicity.  3. The patient bas a histodogically- or cytologically-confirmed diagnosis of treast cancer.  4. The patient beast cancer has halt receptor analysis performed and this is regardive for all the following: the HER2 receptor, oestrogen receptor and progesterone receptor i.e. the patient has the typic negative disease.  5. The patient beast cancer has halt receptor analysis performed and this is regardive of all the following: the HER2 receptor, oestrogen receptor and progesterone receptor i.e. the patient has the typic negative disease.  5. The patient beast cancer has halt receptor analysis performed and this is regardive disease.  7. The patient is dead as being at high risk of recurrences as defined by having T1s N1-2 of 24 asses.  Please indicate below the staging of the breast cancer in this patient:  17. N1-23 disease or  1	No	TA851		_

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PEMB22	Pembrolizumab in combination with chemotherapy with or without bevacizumab	For the treatment of persistent, recurrent or metastatic cervical cancer in patients whose tumour PP-L1 expression test results have a combined positive score (CPS) of 1 or more where the following criteria have been met:	1. This application is being made by and the first cycle of youternic anti-cincer therapy. 2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, collisis, nephroits, entertrologophics, legislation and shall notice. 3. The patient has a bisologically or cyclogically confirmed diagnosis of cervical carcinoma. 3. The patient has a bisologically or cyclogically confirmed diagnosis of cervical carcinoma. 3. The patient has a bisologically or cyclogically confirmed diagnosis of cervical carcinoma. 3. The patient is been bested by an approved and validated test for PD-L1 expression as measured by the combined positive score (CPS) test and the result is 3 or more. Proteins discourse that an article of the patient is presented by the combined positive score (CPS) test and the result is 3 or more. Proteins discourse that all the patient has persistent locoregional discourse with or without distant metastates or has presented with distant metastates or presented with distant metastates or presented to the patient is presented with distant metastates or presented to conglored discourse with or without distant metastates or has presented with distant metastates or presented to conglored discourse with or without distant metastates or has presented with distant metastates or presented to conglored discourse with or without distant metastates or has presented with distant metastates or presented to conglored discourse with or without distant metastates or presented in conglored discourse with distant metastates or presented in conglored discourse with distant metastates or presented in conglored discourse with distant metastates or presented in conglored discourse with distant metastates or presented in conglored discourse with distant metastates or presented in conglored discourse with distant metastates or presented in conglored discourse with distant metastates or presente	No	TA939	13-Dec-23	12-Mar-24

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PEMB23	Pembrolizumab in combination with lenvatinib	For the treatment of patients with endometrial carcinoma who have progressive disease during or following prior platinum-containing therapy given in any setting for advanced or recurrent or metastatic disease and who are not candidates for potentially curative surgery or radiotherapy or themoradiotherapy where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy. 2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinogathies, hepatists and skin toxicity. 3. The patient has a histologically- or cytologically confirmed diagnosis of endometrial carcinoma. Note: patients with endometrial surround of any kind or with carcinoparacoma (Mixed Mullerian tumour) are NOT eligible for pembrolizumab plus lenvatinib. 4. The mismatch repair status of the endometrial carcinoma if known at present: —instracts repair deficient —instracts repair status on the own at present —instracts repair status on the own at present —instracts repair status not known at present —instracts desired and experiment of the patients of the patients repaired and experiment of the patients repaired and experiment of the patients repaired and experiment of the patients repaired and experiment of the patients repaired and experiment of the patients are repaired and experiment of the patient patients are repeated as present patients and experiment of the patients are repaired and experiment of the patients are repaired and experiment of the patients are repaired and experiment of the patients are patients and experiment and include the patients are patients an	No	TA904	21-Jun-23	19-Sep-23

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PEMB24	Pembrolizumab monotherapy	For the subsequent treatment of patients with previously treated unresectable or metastatic COMBECTAL cancer exhibiting microsatellite instability-high (MSI-H) or mismatch repair deficiency (dMMR) when the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy will be prescribed by a consultant specialist specifically trained and accordidated in the use of systemic anti-cancer therapy.  2. The prescribing clinical is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, benefits and skin toxicity.  3. The patient has unresectable or metastatic colorectal carcinoma.  4. The patient's tumour has a documented presence of microstatellite instability-high (MS+H) or DNA mismatch repair deficiency (dMMR) confirmed by validated testing.  5. Wild type or mutant BAS status has been determined on this patient's tumour and the result is recorded below:  - wild type RAS status  - mutant BAS status has been determined on this patient's tumour and the result is recorded below:  - wild type or mutant BAS status has been determined on this patient's tumour and the result is recorded below:  - wild type BAS status  - mutant BAS status	No	TA914	20-Sep-23	19-Dec-23

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with pembrolizumab monotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, hepatitis and skin toxicity.  3. The patient has a histologically- or cytologically confirmed diagnosis of endometrial carcinoma.  Note: patients with endometrial sarcoma of any kind or with carcinosarcoma (Mixed Mullerian tumour) are NOT eligible for pembrolizumab monotherapy.  4. The patient's endometrial carcinoma has documented presence of microsatellite instability (MSI-H) or deficient mismatch repair (dMMR) confirmed by validated testing.  5. The patient has advanced or recurrent or metastatic endometrial carcinoma and is not a candidate for any potentially curative treatment with surgery or radiotherapy or chemoradiotherapy.				
PEMB25	Pembrolizumab monotherapy	For the treatment of patients with ENDOMETRIAL carcinoma exhibiting microsatellite instability (MSI-H) or deficient mismatch repair (dMMR) and who have progressive disease during or following prior platinum-containing therapy given in any settling for advanced	6. The patient has received at least 1 prior platinum-containing chemotherapy given in any setting whether this was as neoadjuvant chemotherapy or as adjuvant therapy or as chemoradiotherapy or for recurrent disease or for metastatic disease or for more than one of these settings.  7. The patient has progressive disease during or following the most recent platinum-containing chemotherapy.  8. Pembrolizumab will be given as monotherapy.  Note: pembrolizumab will be given as monotherapy.  9. The patient has NOT received any prior antibody treatment which targets PD-1 or PD-12 or CD137 or OX40 or anti-cytotoxic T-lymphocyte-associated antigen-4 (CTLA-4).	No	TA914	20-Sep-23	19-Dec-23
		or recurrent or metastatic disease and who are not candidates for potentially curative surgery or radiotherapy or chemoradiotherapy where the following criteria have been met:	10. The patient will be treated with a fixed dose of pembrolizumab of either 200mg every 3 weeks or 400mg every 6 weeks.  Note: NHS England recommends the use of 6-weekly pembrolizumab whenever appropriate.  11. Treatment with pembrolizumab will be stopped at whichever of the following events occurs first: disease progression or loss of clinical benefit or unacceptable toxicity or withdrawal of patient consent or after 2 years of treatment (or after 3 x 3-weekly yedes or its equivalent if 6-weekly pembrolizumab is used).			20-sep-23	19-Dec-23
		12. The patient has an ECOG performance status (PS) of 0 or 1.  Note: NHS England does not fund this treatment in patients of ECOG PS 2.  13. The patient has no symptomatically active brain metastases or leptomeningeal metastases.  14. A formal medical review as to how pembrolizumab is being tolerated and whether treatment should continue or not will be scheduled to occur at least by the end of the second month of treatment.  15. When a treatment break of more than 12 weeks beyond the expected 3 or 6 weekly cycle length is needed, a treatment break approval form will be completed to restart treatment.					
			16. Pembrolizumab will be otherwise used as set out in its Summary of Product Characteristics (SPC).  1. This application is being made by and the first cycle of systemic anti-cancer therapy with pembrolizumab monotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  1. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis,				
		For the subsequent treatment of patients	2. The patient's tumour has a documented presence of microsatellite instability-high (MSI-H) or DNA mismatch repair deficiency (dMMR) confirmed by validated testing.  3. The patient's tumour has a documented presence of microsatellite instability-high (MSI-H) or DNA mismatch repair deficiency (dMMR) confirmed by validated testing.  5. The patient has progressive disease during or following the most recent chemotherapy.				
PEMB26	Pembrolizumab monotherapy	with previously treated unresectable or metastatic <b>GASTRIC</b> cancer exhibiting microsatellite instability-high (MSI-H) or mismatch repair deficiency (dMMR) where the following criteria have been met:	7. The patient has an ECOG performance status (PS) of or 1.  Note: NHS England does not fund this treatment in patients of ECOG PS 2.  8. The patient has no symptomatically active brain metastases or leptomeningeal metastases.  9. The patient has NOT received prior treatment with an anti-PD-1, anti-PD-12, anti-PD-12, or anti-Cytotoxic T-lymphocyte-associated antigen-4 (CTLA-4) antibody.  10. Pembrolizumab will be administered as monotherapy at a dose of 200mg every 3 weeks or a dose of 400mg every 6 weeks.	No	TA914	20-Sep-23	19-Dec-23
		N 11 (o 11 13	Note: NHS England recommends the use of 6-weekly pembrolizumab whenever appropriate.  11. Pembrolizumab will be stopped at whichever of the following events occurs first: disease progression or loss of clinical benefit or unacceptable toxicity or withdrawal of patient consent or after a total treatment duration of 2 years (or a maximum of 35 -weekly cycles or the equivalent number of 6-weekly cycles to result in a total treatment duration of 2 years).  12. A formal medical review as to whether treatment with pembrolizumab should continue will occur at least by the end of the 2nd month of treatment.  13. When a treatment break of more than 12 weeks beyond the expected 3 or 6-weekly cycle length is needed, a treatment break approval form will be completed to restart treatment.  14. Pembrolizumab will be otherwise used as set out in its Summary of Product Characteristics (SPC).				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with pembrolizumab monotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, hepatitis and skin toxicity.				
			3. The patient has unresectable or metastatic small intestinal carcinoma.				
			4. The patient's tumour has a documented presence of microsatellite instability-high (MSI-H) or DNA mismatch repair deficiency (dMMR) confirmed by validated testing.				
			5. The patient has received previous treatment for unresectable or metastatic small intestinal cancer.				
		For the subsequent treatment of patients	6. The patient has progressive disease during or following the most recent chemotherapy.				
PEMB27	Pembrolizumab	with previously treated unresectable or metastatic <b>SMALL INTESTINAL</b> carcinoma exhibiting microsatellite instability-high	7. The patient has an ECOG performance status (PS) of 0 or 1.  Note: NHS England does not fund this treatment in patients of ECOG PS 2.	No	TA914	20-Sep-23	19-Dec-23
PEIVIB2/	monotherapy	(MSI-H) or mismatch repair deficiency	8. The patient has no symptomatically active brain metastases or leptomeningeal metastases.	. NO	1A914	20-Sep-23	19-Dec-23
			9. The patient has NOT received prior treatment with an anti-PD-1, anti-PD-12, anti-CD137, or anti-Cytotoxic T-lymphocyte-associated antigen-4 (CTLA-4) antibody.	1			
		been met:	10. Pembrolizumab will be administered as monotherapy at a dose of 200mg every 3 weeks or a dose of 400mg every 6 weeks.	s			
			Note: NHS England recommends the use of 6-weekly pembrolizumab whenever appropriate.				
			11. Pembrolizumab will be stopped at whichever of the following events occurs first: disease progression or loss of clinical benefit or unacceptable toxicity or withdrawal of patient consent or after a total treatment duration of 2 years (or a maximum of 35 3-weekly cycles or the equivalent number of 6-weekly cycles to result in a total treatment duration of 2 years).				
			12. A formal medical review as to whether treatment with pembrolizumab should continue will occur at least by the end of the 2nd month of treatment.				
			13. When a treatment break of more than 12 weeks beyond the expected 3 or 6-weekly cycle length is needed, a treatment break approval form will be completed to restart treatment.				
			<ol> <li>Pembrolizumab will be otherwise used as set out in its Summary of Product Characteristics (SPC).</li> </ol>				
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with pembrolizumab monotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-L1 treatments including pneumonitis, colitis, nephritis, endocrinopathies, hepatitis and skin toxicity.				
			3. The patient has unresectable or metastatic biliary tract carcinoma.				
			4. The patient's tumour has a documented presence of microsatellite instability-high (MSI-H) or DNA mismatch repair deficiency (dMMR) confirmed by validated testing.				
			5. The patient has received previous chemotherapy for unresectable or metastatic biliary tract cancer.	1			
		For the subsequent treatment of patients	6. The patient has progressive disease during or following the most recent chemotherapy.	1			
		with previously treated unresectable or metastatic BILIARY TRACT cancer	7. The patient has an ECOG performance status (PS) of 0 or 1.	1			
PEMB28	Pembrolizumab	exhibiting microsatellite instability-high	Note: NHS England does not fund this treatment in patients of ECOG PS 2.	No	TA914	20-Sep-23	19-Dec-23
	monotherapy	(MSI-H) or mismatch repair deficiency	8. The patient has no symptomatically active brain metastases or leptomeningeal metastases.			·	
		(dMMR) where the following criteria have	9. The patient has NOT received prior treatment with an anti-PD-1, anti-PD-12, anti-CD137, or anti-Cytotoxic T-lymphocyte-associated antigen-4 (CTLA-4) antibody.				
		been met:	10. Pembrolizumab will be administered as monotherapy at a dose of 200mg every 3 weeks or a dose of 400mg every 6 weeks.				
			Note: NHS England recommends the use of 6-weekly pembrolizumab whenever appropriate.				
			11. Pembrolizumab will be stopped at whichever of the following events occurs first: disease progression or loss of clinical benefit or unacceptable toxicity or withdrawal of patient consent or after a total treatment duration of 2 years (or a maximum of 35 3-weekly cycles or the equivalent number of 6-weekly cycles to result in a total treatment duration of 2 years).				
			12. A formal medical review as to whether treatment with pembrolizumab should continue will occur at least by the end of the 2nd month of treatment.	]			
			13. When a treatment break of more than 12 weeks beyond the expected 3 or 6-weekly cycle length is needed, a treatment break approval form will be completed to restart treatment.	1			
			14. Pembrolizumab will be otherwise used as set out in its Summary of Product Characteristics (SPC).	1			

ueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with pembrolizumab plus chemotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The prescribing clinician is fully aware of the management of and the treatment modifications that may be required for immune-related adverse reactions due to anti-PD-1 or anti-PD-11 treatments including pneumonitis, colitis, nephritis, endocrinopathies, hepatitis and skin toxicity.				
			3. The patient has a histologically- or cytologically-confirmed diagnosis of HER-2 negative adenocarcinoma of the gastro-oesophageal junction or stomach.  Please mark below which site of the primary tumour applies to this patient:  - HER-2 negative adenocarcinoma of the gastro-oesophageal junction  - HER-2 negative adenocarcinoma of the stomach				
			<ol> <li>The patient has locally advanced unresectable or metastatic disease.</li> <li>An approved and validated test has demonstrated that the tumour has a PD-L1 expression with a combined positive score (CPS) of ≥1.</li> </ol>				
			Please document the actual PD-L1 combined positive score (CPS) below: PD-L1 CPS:				
			6. The patient has not received any previous systemic therapy for locally advanced unresectable or metastatic disease i.e. that pembrolizumab plus chemotherapy will be 1st line systemic therapy for locally advanced unresectable or metastatic disease.				
			In addition, please mark below whether the patient has/has not previously received any systemic therapy for earlier stage disease:  - this patient has not received any previous systemic therapy for adenocarcinoma of the gastro-oesophageal junction or stomach  - this patient was previously treated with neoadjuvant chemotherapy for HER-2 negative adenocarcinoma of the gastro-oesophageal junction or stomach and underwent surgery and has since had disease progression  - this patient was previously treated with adjuvant chemotherapy for HER-2 negative adenocarcinoma of the gastro-oesophageal junction or stomach and has since had disease progression  - this patient was previously treated with concurrent chemo-radiotherapy for HER-2 negative adenocarcinoma of the gastro-oesophageal junction with or without surgery and has since had disease progression				
		Pembrolizumab in combination with platinum and fluoropyrimidine-based	7. The patient has not received prior treatment with any antibody which targets PD-1 or PD-L1 or PD-L2 or CD137 or OX40 or anti-cytotoxic T-lymphocyte-associated antigen-4 (CTLA-4) unless the patient discontinued or completed checkpoint inhibitor immunotherapy as part of adjuvant therapy without disease progression and at least 6 months has elapsed between the date of the last immunotherapy treatment and the date of first diagnosis of relapse with recurrent or metastatic disease.				
PEMB29	Pembrolizumab	chemotherapy for previously untreated advanced HER-2 negative gastric or gastro- oesophageal junction adenocarcinoma either of which expresses PD-L1 with a combined positive score of 1 or more where the following criteria have been	Please mark below the appropriate scenario for this patient  - this patient has not received any previous immunotherapy for adenocarcinoma of the gastro-oesophageal junction or stomach  - this patient was previously treated with neoadjuvant platinum-based chemoradiotherapy for adenocarcinoma of the gastro-oesophageal junction and underwent surgery followed by adjuvant nivolumab (NICE TA 713) then discontinued or completed treatment with adjuvant nivolumab without disease progression and this was at least 6 months prior to the first diagnosis of relapse. Please document in the box below the time gap in months between completion of previous adjuvant nivolumab immunotherapy and first diagnosis of disease relapse:	No	TA997	29-Aug-24	27-Nov-2
		met:	Note: the mandatory interval between the last date of administration of any prior adjuvant immunotherapy and first relapse is at least 6 months. For patients suffering a first relapse within 6-12 months of previous immunotherapy, clinicians should bear in mind the long elimination half-lives of immunotherapies and make individual assessments of the overall benefit/risk ratio of re-treatment with immunotherapy.				
			8. The patient has an ECOG performance status (PS) of 0 or 1 and is fit for platinum and fluoropyrimidine-based chemotherapy in combination with pembrolizumab. 9. The patient has no symptomatically active brain metastases or leptomeningeal metastases.				
			10. Pembrolizumab will be administered at a dose of either 200mg 3-weekly or 400mg 6-weekly initially in combination with platinum and fluoropyrimidine-based chemotherapy and subsequently as monotherapy.				
			11. The chemotherapy used in combination with pembrolizumab will be both platinum and fluoropyrimidine-based.  Please mark below which chemotherapy regimen is being used in this patient:				
			- oxaliplatin plus capecitabine - oxaliplatin plus modified de Gramont regimen - cisplatin plus repecitabine - cisplatin plus infused 5-fluorouracil				
			another regimen  12. Pembrolizumab will be stopped at whichever of the following events occurs first: disease progression or unacceptable toxicity or withdrawal of patient consent or after 2 years of treatment (or after 35 x 3-weekly cycles or its equivalent if 6-weekly dosing is used).				
			Note: the 2 year stopping rule for pembrolizumab in this indication was a key part of the company submission to NICE of the clinical and cost effectiveness of pembrolizumab in this indication.  Note: once pembrolizumab is stopped after 2 years of treatment, it cannot be re-started.				
			13. A formal medical review as to how pembrolizumab plus chemotherapy is being tolerated and whether pembrolizumab should continue or not will be scheduled to occur at least by the end of the second 3-weekly cycle of treatment.				
			14. When a treatment break of more than 3 months beyond the expected 3- or 6-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment.  15. Pembrolizumab will otherwise be used as set out in its Summary of Product Characteristics (SPC) with the exception of criterion 12.				

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lueteq Form ref	f: Drug NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
РЕМВЗО	Pembrolizumab in combination with chemotherapy for neoadjuvant treat and then continued as a dijuvant monotherapy in adults with previou untreated UICC/AICC 8th edition stage or IB or III or N2 only III is non-smal lung cancer AND who are candidate potentially curative surgery where following criteria have been met	10. The chemotherapy partner to this neoadjuvant combination will be a 2-drug platinum-based combination with the platinum component being either cisplatin or carboplatin given at a dose of at least AUC of Smg/ml/min.    10   The chemotherapy partner to this neoadjuvant combination will be a 2-drug platinum-based combination with the platinum component being either cisplatin or carboplatin given at a dose of at least AUC of Smg/ml/min.    2   Please mark below which will be the platinum-based component of the 2-drug combination:   3   Composition of the comp	Yes	TA1017	Guidance  20-Nov-24	

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PEMB31	Pembrolizumab monotherapy	Pembrolizumab monotherapy for adjuvant treatment after complete tumour resection in adult patients with UICC/AICC 8th edition stage IIA or IIB or IIIA or N2 only IIIB non-small cell lung cancer and whose disease has not progressed on recently completed adjuvant platinumbased chemotherapy where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with adjuvant pershorolizumab will be personhed by a consustant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The principle control of the properties of the prope	No	TA1037	05-Feb-25	06-May-25

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			13. The patient has not received prior treatment with an anti-PD-1, anti-PD-1, anti-PD-12, anti-PD-137, or anti-Cytotoxic T-lymphocyte-associated antigen-4 (CTLA-4) antibody.				
		Pembrolizumab monotherapy for adjuvant treatment after complete tumour resection in adult patients with UICC/AJCC	14. The patient has not received any neoadjuvant chemotherapy for this NSCLC or any prior or planned adjuvant radiotherapy.				1
			15. The patient has an ECOG performance status (PS) of 0 or 1.				1
	Pembrolizumab	8th edition stage IIA or IIB or IIIA or N2	16. Pembrolizumab will be stopped at whichever of the following events occurs first: disease progression or unacceptable toxicity or withdrawal of patient consent or no completion of 1 year in total duration of treatment with pembrolizumab (i.e. after a maximum of 18 x 3-weekly or 9 x 6-weekly cycles).				ĺ l
PEMB31	monotherapy	only IIIB non-small cell lung cancer and whose disease has not progressed on	penutrukuniau (i.e. aiter a inakinisti en 13.6 3-weeksy Lycies).  17. Penbrolizania wille a diskinistered as monotherapy.	No	TA1037	05-Feb-25	06-May-25
			18. A formal medical review as to how pembrolizumab is being tolerated and whether treatment with pembrolizumab should continue or not will be scheduled to occur at least by the end of the second month of treatment.				1
		based chemotherapy where the following	19. When a treatment break of more than 3 months beyond the expected 3- or 6-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment.	1			
		Criteria have been met:	20. Pembrolizumab will be otherwise used as set out in its Summary of Product Characteristics (SPC).				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PEMB32	Pembrolizumab in combination with platinum-containing chemotherapy (carboplatin and paclitaxel)	Pembrolizumab in combination with platinum-containing chemotherapy (carboplatin and paclitaxel) for the 1st line treatment of mismatch repair deficient (dMMR) or microsatellite instability-high endometrial carcinoma in adult patient who have recurrent or primary advanced disease and who are not candidates for potentially curative surgery or radiotherapy or chemoradiotherapy but are eligible for systemic therapy where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with permission in combination with carboplatin and pacitizate will be prescribed by a consultant specialist specifically trained and accordited in the use of systemic anti-cancer therapy.  2. The patient has an instalogically—or cytologically confirmed diagnosis of endometrial carcinoma (including clear cell and serous histologically—or protogically confirmed diagnosis of endometrial carcinoma (including clear cell and serous histologically—or protogically confirmed deplication).  3. The patient's there has a last recurrence of mismatch repair deficiency (didMMI) or microsatellate instability (MSH-1) confirmed by validated testing.  4. The patient either has a 1st recurrence of endometrial carcinoma after surgery or adoltherapy or chemoradiotherapy or has presented with primary locally advanced or metastatic endometrial carcinoma agal in whichever scenario sign as andiated for any potentially cancellate treatment with surgery or adoltherapy or chemoradiotherapy or chemoradiotherapy or the protogological protogo	No	TA1092	27-Aug-25	25-Nov-25

Blueteq Form ref	: Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PEMB33	Pembrolizumab in combination with platinum-containing chemotherapy (carboplatin and paclitaxel)	Pembrolizumab in combination with platinum-containing chemotherapy (carboplatin and paclitaxel) for the 1st line treatment of mismatch repair proficient (pMMR) or microsatellite stable endometrial carcinoma in adult patients who have recurrent or primary advanced disease and who are not candidates for potentially curative surgery or radiotherapy out are eligible for systemic therapy where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with pemberolizumab in combination with carboplatin and pacifisated will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patients has a thorogocally occupied with combination of the strained of the patients of the strained of th	No	TA1092	27-Aug-25	25-Nov-25

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PEMIG1	Pemigatinib	For locally advanced or metastatic cholangiocarcinoma which has a fibroblas growth factor receptor 2 gene fusion/rearrangement in patients with disease progression during or after previous systemic therapy where the following criteria have been met:	1. This application for pemigatinib is being made by and the first cycle of systemic anti-cancer therapy with pemigatinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a histologically or cytologically confirmed diagnosis of cholangiocarcinoma.  Please also indicate below whether the cholangiocarcinoma is of intrahepatic origin  - the cholangiocarcinoma is of intrahepatic origin  - the cholangiocarcinoma is of extrahepatic origin  - The patient has unresectable locally advanced or metastatic disease.  - The patient has unresectable locally advanced or metastatic disease.  - The patient has been previously treated with systemic therapy for cholangiocarcinoma and the disease has progressed during or after such therapy.  Please also indicate whether the patient has received 1 or 22 lines of systemic therapy for cholangiocarcinoma and the disease has progressed during or after such therapy.  - The patient has been previously treated with systemic therapy for cholangiocarcinoma or the patient has been previously treated with 2 lines of systemic therapy for cholangiocarcinoma  - The patient has been previously treated with 2 lines of systemic therapy for cholangiocarcinoma  - The patient has not previously received any specifically FGFR2-targeted therapy unless furthatinib montherapy has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of progressive disease.  - Please mark below which scenario applies to this patient:  - the patient has not been previously treated with a FGFR2-targeted therapy or furthatinib monotherapy has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of progressive disease.  - The patient has not been previously treated with a FGFR2-targeted therapy or furthatinib monotherapy has had to be stopped within 3 months of its start solely as a consequence of	No	TA722	25-Aug-21	24-Sep-21

26-Nov-2025

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PERZa	Pertuzumab	Neoadjuvant pertuzumab plus trastuzumab in NODE POSITIVE patients for the neoadjuvant treatment of locally advanced, inflammatory or early breast cancer at high risk of recurrence (PER2a) where the following criteria have been met  This form (introduced in November 2019) is for patients known to be pathologically node positive prior to commencing neo-adjuvant therapy. On commencing adjuvant treatment with pertuzumab, form PER4a (for node positive patients) must be completed.  For patients with locally advanced, inflammatory or early breast cancer who are node negative or of unknown nodal status when commencing neoadjuvant pertuzumab, form PER2b must be used for the neoadjuvant part of treatment followed by form PER4b for the adjuvant part of treatment only if the histology post-surgey; is node eve.	1. This application has been made by and the first cycle of systemic anti-cancer therapy.  NOTE: This application should be made immediately prior to commencing pertuzumab plus trastuzumab when given with single agent docetaxel/pacitiaxel chemotherapy as part of sequential anthracycline/taxane regimen and not at the start of the anthracycline based component.  2. Treatment is being initiated with neoadjuvant intent  3. The patient has newly diagnosed locally advanced, inflammatory or early breast cancer at high risk of recurrence (i.e must have stage T2-T4b and M0 disease) and has pathologically-proven node positive disease  4. The patient has HER2 3+ by IHC or FISH/CISH positive disease  5. The patient has a baseline LVEF greater than or equal to 55% % or if anthracyclines were given that the LVEF was greater than or equal to 50% after completion of the anthracycline component of the neo-adjuvant chemotherapy.  6. The patient has received no prior treatment with chemotherapy or HER2 therapy for this breast cancer  7. Perturumab plus trasturumab will be given in combination with docetaxel/pacitiaxel-containing chemotherapy. The exceptions to this are for patients enrolled in the NIHR-approved ROSCO trial (LKCRN Study ID:19059 where neoadjuvant perturumab can be given with chemotherapy in either arm of the study) or potential participants in the NIHR-approved HER2 RADICAL trial (LKCRN Study ID:131362 where pacitiaxel/nab-pacitiaxel/docetaxel may be used). Please indicate below if the patient is enrolled in the NIHR-approved ROSCO or HER2 RADICAL trials  8. The patient will receive a maximum of 4 cycles of perturumab plus trasturumab figuren with single agent docetaxel chemotherapy as part of sequential anthracycline/docetaxel regimen OR 4 cycles of perturumab plus trasturumab if given with swelly pacitiaxels chemotherapy as part of sequential anthracycline/docetaxel regimen OR 4 cycles of perturumab plus trasturumab if given with weekly pacitiaxels chemotherapy as part of sequential anthracycline/docetaxel regim	No	TA424	21-Dec-16	21-Mar-17
			9. Treatment will be given using either intravenous pertuzumab and intravenous biosimilar trastuzumab or using the PHESGO® brand combination pertuzumab and trastuzumab subcutaneous injection.  Please mark as to which mode of administration is to be used:  - Intravenous pertuzumab and intravenous best value biosimilar trastuzumab or  - PHESGO® subcutaneous pertuzumab and trastuzumab combination injection  9. The prescribing clinician understands the differing dosages to be used for the different formulations of pertuzumab and trastuzumab in relation to the first (loading) cycle and then in subsequent cycles:  - Intravenous pertuzumab is given at an initial loading dose of 8 mg/kg body weight followed every 3 weeks thereafter by a maintenance dose of 6 mg/kg body weight				
			**Subcutaneous PHESGO** is given at an initial loading dose of 1,200mg pertuzumab and 600mg trastuzumab in 15 mL of solution in a single-dose vial followed every 3 weeks thereafter by a maintenance dose of 600mg pertuzumab and 600mg trastuzumab in 15 mL of solution in a single-dose vial followed every 3 weeks thereafter by a maintenance dose of 600mg pertuzumab and 600mg trastuzumab in a 10 mL of solution in a single-dose vial.				
			11. Pertuzumab or PHESGO* will be otherwise used as set out in their respective Summary of Product Characteristics (SPCs).				L

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PER2b	Pertuzumab	Neoadjuvant pertuzumab plus trastuzumab in patients who are NODE NEGATIVE or of UNKNOWN NODAL STATUS for the neoadjuvant treatment of locally advanced, inflammatory or early breast cancer at high risk of recurrence (PERZb) where the following criteria have been met:  This form (introduced November 2019) is for patients who are node negative or of unknown nodal status prior to commercing neo adjuvant therapy. If a biopsy post-aurgery shows the patients are found to be node positive, then for them to commence adjuvant treatment with perturumab and ratsuturnab, form PERAD must be completed.  For patients with locally advanced, inflammatory or early breast cancer who are node positive when commencing neo-adjuvant chemotherapy in combination with perturumab and trastuzumab, form PERAD must be used followed by form PERAD when commencing adjuvant treatment with perturumab and trastuzumab.	B. The patient will receive a maximum of 4 cycles of pertuzumab plus trastuzumab if given with single agent docetaxel chemotherapy as part of sequential anthracycline/docetaxel regimen OR 4 cycles of pertuzumab plus trastuzumab if given with weekly paclitaxel chemotherapy as part of sequential anthracycline-paclitaxel regimen OR a maximum of 6 cycles of pertuzumab plus trastuzumab only if given with combination of docetaxel and carboplatin chemotherapy OR a maximum of 4 cycles of pertuzumab plus trastuzumab in given with the first 4 cycles of chemotherapy in either arm of the NIHR-approved ROSCO necadjuvant trial OR a maximum of 6 cycles (minimum of 4) of pertuzumab plus trastuzumab with non-anthracycline taxane containing chemotherapy as part of the NIHR-approved HER2 RADICAL trial of tailored treatment for HER2 positive early breast cancer.  Please indicate below the maximum number of cycles of pertuzumab it is planned for the patient to receive:  - 4 cycles OR  - 6 cycles OR  - Patient is a potential participant in on the HER2 RADICAL necadjuvant trial (4-6 cycles)	No	TA424	21-Dec-16	21-Mar-17

lueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application for pertuzumab in combination with trastuzumab and a taxane or capecitabine is being made by and the first cycle will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			2. The patient has histologically documented breast cancer which is HER2 3+ by immunohistochemistry and/or has a HER2 ratio of ≥2.0 by in situ hybridisation.	1			
			3. The patient has been diagnosed with locally advanced or metastatic breast cancer.	1			
			4. The patient has an ECOG performance status of 0 or 1.				
			5. The patient has a baseline LVEF of greater than or equal to 50%.	1			
			6. Any adjuvant HER2 therapy was completed more than 12 months prior to the diagnosis of locally advanced or metastatic disease.	1			
			7. The patient has had no prior treatment with chemotherapy or HER2 therapy for locally advanced or metastatic disease.				
			8. The patient will receive pertuzumab and trastuzumab as first line treatment in combination with a taxane or capecitabine.				
			9. The prescribing clinican understands that pertuzumab and trastuzumab are not to be used beyond first disease progression outside the CNS.	1			
	Pertuzumab	The first live to start of leastly	Note: Treatment with pertuzumab and trastuzumab can continue if there is disease progression solely within the CNS.				
PER1	(in combination with	The first line treatment of locally advanced or metastatic breast cancer	10. Treatment will be given using either intravenous pertuzumab and intravenous biosimilar trastuzumab or using the PHESGO* brand combination pertuzumab and trastuzumab subcutaneous injection.	Yes	TA509	07-Mar-18	05-Jun-18
	trastuzumab and a taxane	where all the following criteria are met:	Please mark as to which mode of administration is to be used:			07 11101 20	05 7411 20
	or capecitabine)		- Intravenous pertuzumab and intravenous best value biosimilar trastuzumab <b>or</b>				
			- PHESGO® subcutaneous pertuzumab and trastuzumab combination injection				
			11. The prescribing clinician understands the differing dosages to be used for the different formulations of pertuzumab and trastuzumab in relation to the first (loading) cycle and then in subsequent cycles:				
			- Intravenous pertuzumab is given at an initial loading dose of 840mg followed every 3 weeks thereafter by a maintenance dose of 420mg Intravenous trastuzumab is given as an initial loading dose of 8 mg/kg body weight				
			Subcutaneous PHESGO* is given at an initial loading dose of 1,200mg pertuzumab and 600mg trastuzumab and 600mg to 3 weeks thereafter by a maintenance dose will followed every 3 weeks thereafter by a maintenance dose of 6,00mg pertuzumab and 600mg trastuzumab and 6				
			600mg trastuzumab in a 10 mL of solution in a single-dose vial.				
			12. When a treatment break of more than 6 weeks beyond the expected cycle length is needed, I will complete a treatment break form to restart treatment.				
			13. Pertuzumab or PHESGO® will be otherwise used as set out in their respective Summary of Product Characteristics (SPC)				
			1. This application for pertuzumab in combination with trastuzumab as part of adjuvant systemic therapy is made by and the first cycle of adjuvant pertuzumab and trastuzumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			2. The patient has histologically documented breast cancer which is HER2 3+ by immunohistochemistry and/or has a ratio of ≥2.0 by in situ hybridisation.	-			
			3. The patient has been diagnosed with early breast cancer and this has been adequately excised.	-			
			3. The patient has pathologically confirmed suitary lymph node involvement in successful and the pathologically confirmed suitary lymph node involvement.	1			
			Pertuzumab in combination with trastuzumab as adjuvant treatment is only NICE-recommended and commissioned in patients with pathologically documented axillary lymph node involvement.				
			5. The patient is due to commence adjuvant chemotherapy in combination with pertuzumab and trastuzumab and will receive one of the standard adjuvant anthracycline- and/or taxane-based chemotherapy regimens as set out in				
		Pertuzumab in combination with trastuzumab and chemotherapy as	section 4.2 and 5.1 of pertuzumab's Summary of Product Characteristics. Please mark as to which regimen is to be used:				
		adjuvant therapy for axillary node positive	3-3 cycles of FEC or FAC followed by 3-4 cycles of docetaxel or 12 cycles of weekly paclitaxel or 3-4 cycles of AC or EC followed by 3-4 cycles of docetaxel or 12 cycles of weekly paclitaxel or 3-4 cycles of AC or EC followed by 3-4 cycles of ocetaxely or 12 cycles of weekly paclitaxel or				
		HER2-positive early breast cancer and with	1 - 3-4 tytes of AC of EC followed by 3-4 tytes of acceptance of 12 tytes of weeking patriatate of 1-6 tytes of occeptance and acceptance of 12 tytes of weeking patriatate of 1-6 tytes of occeptance and acceptance of 12 tytes of weeking patriatate of 1-3-4 tytes of AC of 12 tytes of AC of 13 tytes o				
		NO preceding neoadjuvant chemotherapy	Pertuzumab and trastuzumab should start following completion of the entire anthracycline regimen if given. Pertuzumab and trastuzumab should commence with the first taxane cycle. Pertuzumab and trastuzumab are not				
		in combination with pertuzumab and	commissioned in combination with other adjuvant chemotherapy regimens.				
		criteria have been met:	If a patient has a severe allergic reaction to the docetaxel part of the treatment combination, the patient can be switched to a trial of weekly paclitaxel.				
			6. A maximum of 18 cycles of pertuzumab plus trastuzumab will be administered as adjuvant treatment.	-			
PER3	Pertuzumab	Note: there is a separate form PER4a for adjuvant pertuzumab for node positive patients who	7. Treatment will be given using either intravenous pertuzumab and intravenous biosimilar trastuzumab or using the PHESGO* brand combination pertuzumab and trastuzumab subcutaneous injection.	No	TA569	20-Mar-19	18-Jun-19
PERS	Pertuzuman	received neoadjuvant chemotherapy in	Please mark as to which mode of administration is to be used:	INO	1A509	20-Mar-19	18-Jun-19
		combination with pertuzumab and trastuzumab and who continue on to adjuvant treatment after	- Intravenous pertuzumab and intravenous best value biosimilar trastuzumab or - PHESGO® subcutaneous pertuzumab and trastuzumab combination injection				
		surgery.	- Pricssor's succutaneous pertuzumao and d'astuzumao combination injection				
		For patients who were node negative or of	8. The prescribing clinician understands the differing dosages to be used for the different formulations of pertuzumab and trastuzumab in relation to the first (loading) cycle and then in subsequent cycles:	-			
		unknown nodal status when commencing neo-	1. Intravenous perturumab is given at an initial loading dose of 840mg followed every 3 weeks thereafter by a maintenance dose of 420mg.				
		adjuvant chemotherapy in combination with pertuzumab and trastuzumab and in whom	- Intravenous trastuzumab is given as an initial loading dose of 8 mg/kg body weight followed every 3 weeks thereafter by a maintenance dose of 6 mg/kg body weight				
		surgery has demonstrated node positive disease,	- Subcutaneous PHESGO* is given at an initial loading dose of 1,200mg pertuzumab and 600mg trastuzumab in 15 mL of solution in a single-dose vial followed every 3 weeks thereafter by a maintenance dose of 600mg pertuzumab and				
		form PER4b must be used for adjuvant	600mg trastuzumab in a 10 mL of solution in a single-dose vial.				
		pertuzumab.		]			
			9. The patient has an ECOG performance status of 0 or 1.	]			
			10. The pre-treatment left ventricular ejection fraction was ≥55% and if anthracyclines were given that the LVEF was ≥50% after completion of the anthracycline component of the adjuvant chemotherapy.				
				]			
			111. When a treatment break of more than 6 weeks beyond the expected cycle length is needed. I will complete a treatment break form to restart treatment, including an indication as appropriate if the nation that an extended break				
			11. When a treatment break of more than 6 weeks beyond the expected cycle length is needed, I will complete a treatment break form to restart treatment, including an indication as appropriate if the patient had an extended break because of COVID 19.				

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PER4a	Pertuzumab	Pertuzumab in combination with trastuzumab as adjuvant therapy for patients with HER2-positive early breast cancer which was diagnosed as being NODE POSITIVE prior to neoadjuvant treatment and has now completed neoadjuvant pertuzumab in combination with trastuzumab and chemotherapy and surgery (PER4a) where the following criteria have been met:  These patients must have had form PER2a completed for the neoadjuvant portion of their therapy.  For patients who were node negative or of unknown nodal status prior to commencing no-adjuvant therapy, form PER2b (neoadjuvant pertuzumab in such PER2b patients who are found to be node positive after surgery.  For node positive patients who did not receive neo-adjuvant therapy with pertuzumab form PER3 should be used for adjuvant treatment of pertuzumab + trastuzumab.	1. This application for pertuzumab in combination with trastuzumab as part of adjuvant chemotherapy is made by and the first cycle of adjuvant pertuzumab and trastuzumab will be prescribed by a consultant specialist specifically trained and accretified in the use of systemic anti-cancer therapy.  2. The patient has histologically documented breast cancer which is HER2 3+ by immunohistochemistry and/or has a ratio of \$2.0 by in situ hybridisation.  3. The patient has been diagnosed with early breast cancer and this has been adequated yexised.  4. Experience in terms of the invasive carcinoma to neadjuvant chemotherapy in combination with perturnab and trasturumab: pathological complete response in horse of the invasive carcinoma to neadjuvant chemotherapy in combination with perturnab and trasturumab or residual invasive disease remaining in breast and aliany nodes after neadjuvant chemotherapy in combination with perturnab and trasturumab - unknown (patient started on adjuvant perturnab post-surgery as they were known to be node positive before the pathology results were available to confirm the status as to pathological complete response in horse of perturnab plus trasturumab post-surgery as they were known to be node positive before the pathology results were available to confirm the status as to pathological complete remission.  5. The patient had confirmed node positive disease prior to neo-adjuvant treatment and surgery  6. A maximum of 18 cycles of perturnab plus trasturumab pust-surgery as they were known to be node positive and before the pathology results have confirmed the status as to pathological complete remission are given in combination with neoadjuvant chemotherapy, then a maximum of 14 cycles of adjuvant perturnab and trasturumab No	TA569	20-Mar-19	18-Jun-19	

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
PER4b	Pertuzumab	Pertuzumab in combination with trastuzumab as adjuvant therapy for HER2 positive early breast cancer patients thought to be node negative or of unknown nodal status prior to neoadjuvant chemotherapy and found to be axiliarly node positive AFTER completion of neoadjuvant pertuzumab/frastuzumab and surgery (PER4b) where the following criteria have been met:  These patients must have completed form PER2b for the neoadjuvant portion of their therapy.  PER2b patients (node negative or of unknown nodal status prior to neoadjuvant chemotherapy) who are node negative after surgery cannot have adjuvant perturumab as NICE has only recommended adjuvant perturumab in patients who are node positive.  For patients known to be node positive prior to commencing neoadjuvant therapy, forms/PER2a (neoadjuvant portion of treatment) and PER4a (adjuvant portion of treatment) must be used.  For node positive patients who did not receive neoadjuvant themotherapy, applications for adjuvant pertuzumab should proceed directly to adjuvant terturumab should proceed directly to adjuvant terturumab should proceed directly to adjuvant terturumab inform PER3).	1. This application for perturumab in combination with tratturumab as part of adjuvant chemotherapy is made by and the first cycle of adjuvant perturumab and trastzumab will be prescribed by a consultant specialist specifically ratined and accredited in the use of systemic anti-cancer threapy.  2. The patient has histologically documented breast cancer which is HER2 3- by immunohistochemistry and/or has a ratio of 22.0 by in situ hybridisation.  3. The patient has been diagnosed with early breast cancer and this has been adequately excised.  4. The patient has received monodipunot chemotherapy in combination with perturumab and trastrurumab or pathological complete response in the breast but not in the avillary nodes after neoadjuvant chemotherapy in combination with perturumab and trastrurumab or pathological complete response in the breast but not in the avillary nodes after neoadjuvant chemotherapy in combination with perturumab and trastrurumab or residual invasive disease remaining in both breast and avillary nodes after neoadjuvant chemotherapy in combination with perturumab and trastrurumab or residual invasive disease remaining in both breast and avillary nodes after neoadjuvant chemotherapy in combination with perturumab and trastrurumab or security of the pathological complete response in the breast but not in the avillary nodes after neoadjuvant treatment and definitive surgery has since found residual invasive carcinoma in the avillary nodes after neoadjuvant treatment and definitive surgery has since found an absence of invasive carcinoma in the avillary nodes but there are histological changes (such as fibrosis) which the pathologist has interpreted as representing previous avillary nodal involvement  5. A maximum of 18 cycles of perturumab public trastrurumab will be administered during the whole treatment period of neoadjuvant treatment addinitive surgery has si	No	TA569	20-Mar-19	18-Jun-19

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lueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is being made by and the first cycle of systemic anti-cancer therapy.  2. The patient is either an adult (age >=18 years) or a post-pubescent child (age <18 years). Please mark below whether the patient is a post-pubescent child:  - the patient is a post-pubescent child*  - the patient is a post-pubescent child*  - the patient is a post-pubescent child*  - the patient is a post-pubescent child*  - the patient is a post-pubescent child*  - the patient is a post-pubescent child*  - Please note the use of polatuzumab vedotin in combination with bendamustine and rituximab is unlicensed in under 18 year old patients so the Trust policy regarding the use of unlicensed medicines should be followed.  3. The patient has a histologically confirmed diagnosis of diffuse large B cell lymphoma (DLBCL). This includes the following:  - DLBCL not otherwise specified (NCS) (including germinal centre B-cell (GCB)and activated B-cell (ABC) subtypes)  - primary mediastinal large B cell lymphoma  - Epstein-Barr virus (EBV) positive DLBCL  - intravascular rage B cell lymphoma  - double hit and triple hit high grade B cell lymphoma  - double hit and triple hit high grade B cell lymphoma  - double hit and triple hit high grade B cell lymphoma  - double hit and triple hit high grade B cell lymphoma  - double hit and triple hit high grade B cell lymphoma  - double hit and triple hit high grade B cell lymphoma  - double hit and triple hit high grade B cell lymphoma  - double hit and triple hit high grade B cell lymphoma  - double hit and triple hit high grade B cell lymphoma  - double hit and triple hit high grade B cell lymphoma  - double hit and triple hit high grade B cell lymphoma  - double hit and triple hit high grade B cell lymphoma (BPC, BCD, BCD, BCD, BCD, BCD, BCD, BCD, BC				started
POL1	Polatuzumab vedotin in combination with bendamustine and rituximab	For previously treated patients with relapsed or refractory diffuse large B-cell lymphoma and who are not candidates for haematopoietic stem cell transplantation where the following criteria have been met:	SCT centre representation. Please record in the box below which of the following best applies to this patient:	No	TA649	23-Sep-20	23-Oct-20
			7. Treatment with polatuzumab vedotin will be used in combination only with bendamustine and the intravenous formulation of rituximab.  8. Either the patient has not been previously treated with bendamustine for DLBCL or if the patient has been treated previously with bendamustine for DLBCL, this application is to continue a previous registration for the polatuzumab EAMS scheme or the interim polatuzumab Covid-19 access or the patient received bendamustine as part of combination treatment with polatuzumab for bridging therapy to CAR-T cell treatment or if treated with bendamustine outside either of these three options, then the response duration to that course of treatment with bendamustine for DLBCL exceeded 1 year.  9. The patient has an ECOG performance status score of 0 or 1 or 2.  10. The patient will be treated with a maximum of six 3-weekly cycles of polatuzumab vedotin in combination with bendamustine and rituximab.  11. The prescribing clinician understands that the use of bendamustine in this DLBCL indication is unlicensed and that Trust policy regarding the use unlicensed treatments has been followed.  12. The prescribing clinician is fully aware of the MHRA warning in July 2017 that increased mortality has been observed in recent clinical studies in off-label use of bendamustine and that patients need to be monitored for opportunistic infection and hepatitis 8 reactivation.  13. A formal medical review as to whether treatment with polatuzumab in combination with bendamustine plus rituximab should continue or not will be scheduled to occur at least by the end of the first 6 weeks of treatment.  14. When a treatment break of more than 6 weeks beyond the expected 3-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment, including as appropriate if the patient had an extended break on account of Covid-19.  15. Polatuzumab vedotin, bendamustine and rituximab will otherwise be used as set out in their respective Summary of Product Characteristics SPCs).				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
POL2_v1.2	Polatuzumab vedotin in combination with rituximab, cyclophosphamide, doxorubicin and prednisolone	For people with previously untreated diffuse large B-cell lymphoma where the following criteria have been met:	LTGs application is being made by and dato the first cycle of presented per control through the control through through the control through through the control through through the control through through through the control through through through the control through th	No	TA874	01-Mar-23	30-May-23

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application for pomalidomide has been made by and the first cycle of systemic anti-cancer therapy with pomalidomide will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy				
			2. The patient has multiple myeloma				
	POM1 Pomalidomide	Pomalidomide for multiple myeloma previously treated with lenalidomide and bortezomib  3. The patient's performance status (PS) is 0-2  4. The patient has previously received 3 lines of treatment with adequate trials of at least all of the following options of therapy: a routinely commissioned or CDF-funded proteasome inhibitor (bortezomib/carfilzomib/ixazomib), lenalidomide and alkylating agents	No				
POM1	Pomalidomide			TA427	11-Jan-17	11-Apr-17	
			5. The patient has refractory disease to the previous line of treatment				
			6. Pomalidomide will be used as outlined in the Summary of Product Characteristics (SPC)				
		The treatment of Philadelphia	1. An application has been made by and the first cycle of systemic anti-cancer therapy is to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy				
PON1	Ponatinib	chromosome positive acute lymphoblastic leukaemia where all the following criteria	2. The patient has Philadelphia chromosome positive acute lymphoblastic leukaemia	Yes	TA451	13-Feb-17	26-Sep-17
		are met:	3. Imatinib is not clinically appropriate for the patient or the T315I gene mutation is present				
		The treatment of chronic phase,	1. An application has been made by and the first cycle of systemic anti-cancer therapy is to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy				
PON6	Ponatinib	accelerated phase or blast phase chronic myeloid leukaemia where all the following	2. The patient has chronic phase, accelerated phase or blast phase chronic myeloid leukaemia	Yes	TA451	13-Feb-17	26-Sep-17
	1 '	myeloid leukaemia where all the following	3. The disease is resistant to dasatinib or nilotinib, or the patient cannot have dasatinib nor nilotinib and imatinib is not clinically appropriate, or the T315I gene mutation is present				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
QUIZ1	Quizartinib	For the treatment of adult patients for treating newly diagnosed FLT3-TD mutation positive acute myeloid leukaemia where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with quizartinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient is an adult and has a confirmed diagnosis of acute myeloid leukaemia.  3. The patient's AML F.T3-ITD mutation as determined by a validated test.  Note: quizartinib is not commissioned for use in patients with AML bearing a FLT3-TKD mutation.  4. The patient is newly diagnosed with FLT3-ITD positive acute myeloid leukaemia and either has not received any induction chemotherapy whilst awaiting FLT3 status.  Please record the status as to induction chemotherapy: - the patient has not vet received any induction chemotherapy or - the patient has received only a single cycle of induction chemotherapy or - the patient has received only a single cycle of induction chemotherapy whilst awaiting the FLT3 result  5. The patient is fit for intensive induction chemotherapy.  6. The patient will be treated with quizartinib only in combination with standard anthracycline and cytarabine induction chemotherapy and then in combination with high dose cytarabine consolidation chemotherapy.  Quizartinib is excluded from the NHS England Treatment Breaks Policy.  7. As maintenance monotherapy, quizartinib is to be only used in patients in complete remission of their AML.  8. In the maintenance monotherapy, quizartinib is to be only used in patients in complete remission of their AML.  8. In the maintenance monotherapy phase, a maximum of 36 x 28-day cycles of quizartinib can be re-started subject to the maximum total maintenance treatment duration of 36 x 28 day cycles.  10. In view of the patient has undergone a stem cell transplant, maintenance quizartinib can be re-started subject to the maximum total maintenance received maintenance evolving trion gradiation chemotherapy, once weekly during the LST amount of a complete remained and consolidation chemotherapy, once weekly during the LST	No	TA1013	23-Oct-24	21-Jan-25

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
N/A	Radium-223	Radium-223 dichloride for treating hormone-relapsed prostate cancer with bone metastases	1. This application has been made by and the first cycle of systemic anti-cancer therapy with radium-223 will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. ONE of the following applies to this patient:  - The patient has histologically or cytologically confirmed adenocarcinoma of the prostate and has castration-resistant disease with two or more symptomatic bone metastases detected on skeletal scintigraphy OR  - The patient has histologically or cytologically cytologically or cytologically or cytologically or cytologically or cytologically or cytologically or cytologically or cytologically or cytologically or cytologically or cytologically or cytologically or cytologically or cytologically or cytologically or cytologically or cytologically or cytologically or cytologically cytologically or cytological	Yes	TA412	28-Sep-16	28-Dec-16
REG1	Regorafenib	The treatment of previously treated unresectable or metastatic gastrointestinal stromal tumours where al the following criteria are met:	1. An application has been made by and the first cycle of systemic anti-cancer therapy is to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. Patient has histologically confirmed, metastatic or unresectable GIST  3. Patient has ECOG performance status (PS) 0-1  4. Patient has had disease progression on or intolerance to previous imatinib  5. Patient has had disease progression on or intolerance to previous sunitinib  6. No treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve)  7. Regorafenib to be otherwise used as set out in its Summary of Product Characteristics	Yes	TA488	15-Nov-17	14-Feb-18

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
REG2_v1.1	Regorafenib	The second line of tyrosine kinase inhibitor systemic therapy of Child-Pugh A locally advanced or metastatic hepatocellular carcinoma previously treated with sorafenib where the following criteria are met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with regorafenib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient turrently has Child-Pugh liver function class A.  Note: NICE has not recommended regorafenib for patients with Child-Pugh liver function class B.  4. The prescribing clinician is aware that there is no efficacy and toxicity data for regorafenib in patients previously treated with sorafenib who had to either discontinue sorafenib on account of toxicity or were unable to tolerate total daily doses of sorafenib of 400mg or more.  5. The patient has an ECOS performance status of 0 or 1.  Note: NICE has not recommended regorafenib in patients with an ECOS performance status of ≥2.  6. The only other TIX with which the patient has been previously treated is sorafenib unless cabozantinib has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression.  7. The patient has not been previously treated with regorafenib.  8. Regorafenib is to be used only as monotherapy.  9. Regorafenib is to be used only as monotherapy.  10. A formal medical review as to whether treatment with regorafenib should continue or not will be scheduled to occur no later than by the end of the 2nd month of therapy.  11. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, a treatment break form will be completed to restart treatment, including an indication as appropriate if the patient had an extended break because of COVID 19.	No	TASSS	09-Jan-19	09-Apr-19
REG3	Regorafenib	For patients with either metastatic or locally advanced and inoperable colorectal cancer who have been previously treated with, or are not considered candidates for, available therapies including fluoropyrimidine-based chemotherapy and anti-EGFR-based treatment where the following criteria have been met:	1. This application is both being made by and the first cycle of systemic anti-cancer therapy with regorafenib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a histologically confirmed diagnosis of adenocarcinoma of the colon or rectum.  3. The patient has netastatic or locally advanced and inoperable disease.  4. The patient has netastatic or locally advanced and inoperable disease.  4. The patient has been previously treated for metastatic disease with, or is not considered a candidate for, fluoropyrimidine-containing chemotherapies which include 5-fluorouracil and/or capecitabine and/or tegafur but not necessarily trifluridine (plus tipiracil).  5. The patient has been previously treated with, or is not considered a candidate for, anti-EGFR-containing chemotherapy.  6. If the patient has been previously breated with trifluridine plus tipiracil (with or without bevacizumab) or not.  Please tick which option applies to this patient:	No	TA866	08-Feb-23	09-May-23

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
RIB1_v1.4	Ribociclib (in combination with an aromatase inhibitor)	The treatment of previously untreated, hormone receptor-positive, HER2-negative, locally advanced or metastatic breast cancer	1. This application for inlocicible in combination with an aromatase inhibitor is made by and the first cycle of systemic anti-cancer therapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The patient has histologically or cytologically documented oestrogen receptor positive and her-2 negative breast cancer  3. The patient has histologically or cytologically documented oestrogen receptor positive and her-2 negative breast cancer  3. The patient has had no prior treatment with a CDK 4/6 inhibitor or locity and in the clear absence of disease progression or a CDK 4/6 inhibitor has been previously received as adjuvant therapy and treatment was completed without disease progression at least 12 months prior to the first diagnosis of recurrent or metastatic disease.  Please mark below which one of these 4 scenarios applies to this patient:  1. no prior treatment with a CDK 4/6 inhibitor or or or previously received as the control of the	No	TA496	20-Dec-17	20-Mar-18
RIB2	Ribociclib in combination with fulvestrant	The treatment of hormone receptor- positive, HER2-negative, locally advanced or metastatic breast cancer where the following criteria have been met:	1. This application for ribocicibl in combination with fulvestrant is being made by and the first cycle of ribocicibl plus fulvestrant will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anticancer therapy.  2. The patient has histologically or cytologically documented oestrogen receptor positive and HER-2 negative breast cancer.  3. The patient has instologically or cytologically advanced breast cancer which is not amenable to curative treatment.  4. The patient is male or is female and if female is either post-menopausal or if pre- or peri-menopausal has undergone ovarian ablation or suppression with LHRH agonist treatment.  5. The patient has an ECOS performance status of 0 or 1 or 2.  6. The patient has an ECOS performance status of 0 or 1 or 2.  6. The patient has seed or seven the patient falls into:  • The patient has an ECOS performance status of 0 or 1 or 2.  6. The patient has seed of previous endocrine therapy according to one of the three populations as set out below as these are the groups on which the NICE Technology Appraisal for ribocicib plus fulvestrant focused. Please record which oppulation the patient falls into:  • The patient has received previous endocrine therapy according to one of the three populations as set out below as these are the groups on which the NICE Technology Appraisal for ribocicib plus fulvestrant focused. Please record which oppulation the patient falls into:  • The patient has been previously exceived following disease progression or  • The patient has been previously disease whilst still receiving adjuvant or neoadjuvant endocrine therapy for early breast cancer with no subsequent endocrine therapy received following disease progression or  • The patient has had no prior treatment with a COK 4/6 inhibitor understatic disease progression or a COK 4/6 inhibitor or does—inhibitor patients and according to the patients of does—inhibitor solicity and in the clear absence of disease progression at least 12 months prior to	No	TA687	31-Mar-21	29-Jun-21

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
RIB3	Ribociclib in combination with an aromatase inhibitor	Ribociclib in combination with an aromatase inhibitor as adjuvant treatment for high risk hormone receptor-positive and HER2-negative early breast cancer where the following criteria have been met:	1. This application for ribocicit bin combination with an aromatase inhibitor is being made by and the first cycle of ribocicit bin plan an aromatase inhibitor will be prescribed by a consultant specialist specifically trained and accredited in the use of yuterian activation and control training.  2. The patient has early becast cancer as defined by having one of the following combinations of 7 and N stage, number of involved availary nodes, histological grade, 1687 index or gene signature.  Please mark in the box below which citegory describes the disease staging of this patient's breast cancer:  1. This grade to or grade 2 disease with 1.3 positive availary nodes or  1. This grade to or grade 2 disease with 1.3 positive availary nodes or  1. This grade to grade 2 disease with 1.3 positive availary nodes or  1. This grade to grade 2 disease with 1.3 positive availary nodes or  1. This grade to grade 2 disease with 1.3 positive availary nodes or  1. This grade to grade 2 disease with 1.3 positive availary nodes or  1. This grade to grade 2 disease with 1.3 positive availary nodes or  1. This grade to grade 2 disease with 1.3 positive availary nodes or  1. This disease of any grade or  1. This disease of any grade or  1. This disease of any grade or  1. This disease of any grade or  1. This disease of any grade or  1. Spositive availary hymph nodes and a primary tumour size 25cm or  1. Spositive availary hymph nodes and a primary tumour size 25cm and histological grade 3 disease  5. The patient has completed any adjuvant or necodiporant themotherapy or  1. Spositive availary hymph nodes and a primary tumour size 25cm and histological grade 3 disease  5. The patient has completed any adjuvant or necodiporant themotherapy or  1. Spositive availary hymph nodes and a primary tumour size 25cm and histological grade 3 disease  5. The patient has completed any adjuvant or necodiporant themotherapy or  1. Spositive availary hymph nodes and a primary tumour size 25cm and histological grade 3 disease  5. The patient has b	Yes	TA1086	06-Aug-25	04-Nov-25

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
RUC1	Rucaparib	As maintenance treatment in patients with high grade epithelial ovarian, fallopian tube or primary peritoneal carcinoma who have a deleterious or suspected deleterious germline and/or somatic BRCA mutation and who have a recent FIRST OR SUBSEQUENT relapse of platinum-essentitive disease and who are now in response following a SECOND OR SUBSEQUENT platinum-based chemotherapy where the following criterihave been met:  There is a separate form (RUC2) for rucaparib as maintenance treatment in patients with high grade epithelial ovarian fallopian tube or primary peritoneal carcinoma who do NOT have deleterious or suspected deleterious germline and/or somatic BRCA mutation and who are in response following platinum-based SECOND or subsequent line chemotherapi	9. The patient is currently less than 8 weeks from the date of the last infusion of the last cycle of the 2nd or subsequent line of platinum-based chemotherapy.  10. The patient has not previously received any PARP inhibitor unless claparil or niraparily but the CDF has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease nonrescription. Please make helow which if the four scenarios analies to this patient:	Yes	TA1007	17-Sep-24	17-Oct-24

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
RUC2	Rucaparib	or suspected deleterious germline and/or somatic BKCA mutation and who have a recent FIRST OR SUBSEQUENT relapse of platinum-sensitive disease and who are now in response following a SECOND OR SUBSEQUENT line platinum-based chemotherapy where the following criteria have been met:  There is a separate form RUC1 for rucaparib as maintenance treatment in patients with high grade epithelial stage lill or IV ovarian, fallopian tube or primary peritoneal carcinoma who have a deleterious or suspected deleterious	6. The patient has recently completed a further line of platinum-based chemotherapy and has received a minimum of 4 cycles of platinum-based treatment. Please enter below what line of platinum-based treatment was the most recent line of treatment:  7. This patient has responded to the recently completed SECOND or subsequent line platinum-based chemotherapy and has achieved a partial or complete response to treatment according to the definitions given below and there is no evidence of progressive disease on the post-treatment scan or a rising CA125 level.  Please enter below as to which response assessment applies to this patient:  - achieved a complete response at the end of the 2nd or subsequent line of platinum-based chemotherapy i.e. has no measurable or non-measurable disease on the post-chemotherapy scan and the CA125 is normal achieved a complete response at the end of the 2nd or subsequent line of platinum-based chemotherapy i.e. has had at least a 30% reduction in measurable or non-measurable disease from the start of to the completion of the 2nd platinum-based chemotherapy or the patient has a complete remission on the post-chemotherapy or T scan but the CA125 has not decreased to within the normal range.  8. The patient is currently less than 8 weeks from the date of the last infusion of the last cycle of the recent 2nd or subsequent line platinum-based chemotherapy.  9. The patient has not previously received any PARP inhibitor or unless either niraparib via the CDF has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression or the patient has previously received any PARP inhibitor or the patient meets all the other criteria listed here.  10. Rucaparib will be used as monotherapy.  11. The patient has an ECO6 performance status of either 0 or 1.	Yes	TA1007	17-Sep-24	17-Oct-24

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			la companya di mangantan di mangantan di mangantan di mangantan di mangantan di mangantan di mangantan di mang	drug/ indication	TA	NICE Guidance	baseline funding started
RUC3	Rucaparib	As maintenance treatment in patients with high grade epithelial stage III or IV ovarian, fallopian tube or primary peritoneal carcinoma who are in response following platinum-based FIRST line chemotherapy AND who DO NOT HAVE deleterious or suspected deleterious BRCA germline and/or somatic BRCA mutation BUT DO HAVE a positive status for homologous recombination deficiency as defined by the presence of genomic instability where the following criteria have been met:	1. This application for maintenance rucapanis is being made by and the first cycle of systemic anticancer therapy with rucapanis will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anticancer therapy with rucapanis will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anticancer therapy with rucapanis will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anticancer therapy will be prescribed by a possible than the germline and/or somatic fumour BRCA testing.  1. This patient has the germline and/or somatic fumour BRCA testing.  1. The patient has testificated and the system of the patient by the possible will be prescribed by a possible five financer or regardly somatic BRCA mutation test with somatic RRCA mutation test will be made the patient by the patient by the possible of the patient by the pat	Yes	TA1055	16-Apr-25	15-Jul-25
			1.1 The patient will commence maintenance rucaparib monotherapy within 8 weeks from the date of the first day of the last cycle of 1st line chemotherapy unless the patient was previously entered into the company's early access scheme for maintenance rucaparib after 1st line chemotherapy and all the other treatment criteria set out in this form are fulfilled.  12. The patient has not previously received any PARP inhibitor unless either the patient has received rucaparib as part of a company early access scheme for this 1st line maintenance indication and the patient meets all the other criteria set out in this form or 1st line maintenance niraparib monotherapy has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression. Such patients must have a positive status for HRD and a negative status for a BRCA mutation.				
			Please mark below which scenario applies to this patient: - the patient has never previously received a PARP inhibitor or - the patient has received rucaparib as part of a company early access scheme for this 1st line maintenance indication and all the other criteria set out in this form are fulfilled - the patient has previously received niraparib monotherapy as 1st line maintenance therapy and this has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression and all the other criteria on this form are fulfilled. By ticking this box, you are confirming that the patient has HRD-positive and BRCA-negative disease.				

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
RUC3 (CONT)	Rucaparib	ovarian, fallopian tube or primary peritoneal carcinoma who are in response following platinum-based FIRST line chemotherapy AND who DO NOT HAVE deleterious or suspected deleterious BRCA germline and/or somatic BRCA mutation BUT DO HAVE a positive status for homologous recombination deficiency as	- the prescribing clinician has discussed with the patient that rucaparib in this indication is less effective than olaparib plus bevacizumab but less costly	Yes	TA1055	16-Apr-25	15-Jul-25

Slueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
RUC4	Rucaparib	As maintenance treatment in patients with high grade epithelial stage III or IV ovarian, fallopian tube or primary peritoneal carcinoma who are in response following platinum-based FIRST line chemotherapy for a tumour which has a NEGATIVE status for a deleterious or suspected deleterious BRCA germline and/or somatic BRCA mutation AND a NEGATIVE or UNKNOWN Status for homologous recombination deficiency as defined by the presence of genomic instability where the following criteria have been met:	L. This application for maintenance rucaparib is being made by and the first cycle of systemic anticiancer therapy.  A. This patient has a proven histological diagnosis of predominantly high grade serous or high grade endometriol of high grade clear cell ovarian, fallopian tube or primary peritoneal carcinoma.  Please enter below as to which is the predominant histology in this patient.  High grade close decoccarcinoma or high grade declaration of the property of the patient of the property of the patients has stage in disease and had an upfront attempt a toptimal cytoreductive surgery and had not visible disease at the end of surgery or the patient has stage in disease and had an upfront attempt a cytoreductive surgery and had not visible disease at the end of surgery or the patient has stage in disease and had an upfront attempt a cytoreductive surgery and had not visible disease	Yes	TA1055	16-Apr-25	started started

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
		As maintenance treatment in patients	10. Maintenance bevacizumab is NOT a treatment option because the patient is not eligible for maintenance bevacizumab monotherapy as set out in form BEV10 or the use of bevacizumab is contraindicated or the maintenance bevacizumab has had to be discontinued within 3 months of its start on account of unacceptable toxicity and in the clear absence of disease progression and all the other criteria on this form are fulfilled.  11. The patient will commence maintenance rucaparib monotherapy within 8 weeks from the date of the first day of the last cycle of 1st line chemotherapy unless the patient was previously entered into the company's early access scheme for maintenance rucaparib after 1st line chemotherapy and all the other treatment criteria set out in this form are fulfilled.				
		with high grade epithelial stage III or IV ovarian, fallopian tube or primary peritoneal carcinoma who are in response following platinum-based FIRST line chemotherapy for a tumour which has a	12. The patient has not previously received any PARP inhibitor unless either the patient has received rucaparib as part of a company early access scheme for this 1st line maintenance indication and the patient meets all the other criteria set out in this form or 1st line maintenance niraparib monotherapy has had to be stopped within 3 months of its start solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression and all the other criteria on this form are fulfilled.  13. Rucaparib will be used as monotherapy.				
RUC4 (CONT)	Rucaparib	NEGATIVE status for a deleterious or suspected deleterious BRCA germline and/or somatic BRCA mutation AND a NEGATIVE or UNKNOWN status for homologous recombination deficiency as	14. Maintenance rucaparib is not being administered concurrently with maintenance bevacizumab.  15. The patient has an ECOG performance status of either 0 or 1.  Note: a patient with a performance status of 2 or more is not eligible for rucaparib.	Yes	TA1055	16-Apr-25	15-Jul-25
	defined by the presence of genomic instability where the following criteria have been met:  Note: NICE's decision as regards the clinical and cost effectiveness of rucaparib in this indication was based on the application of a 2 year calendar year for stopping treatment, i.e. treatment is stopped 2 calendar years after starting, irrespective of treatment breaks.  17. A first formal medical review as to whether maintenance treatment with rucaparib should continue or not will be scheduled to occur at least by the start of the third 4-weekly cycle of treatment.  18. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment.						
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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
RUX1_v2.1	Ruxolitinib	Ruxolitinib for treating disease-related spienomegaly or symptoms in adults with intermediate-2 or high-risk myelofibrosis where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with rusolitinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has primary myelofibrosis (also known as chronic idiopathic myelofibrosis) or post polycythaemia vera myelofibrosis or post essential thrombocythaemia myelofibrosis.  2. The patient has primary myelofibrosis of post polycythaemia vera myelofibrosis or post polycythaemia polycythaemia myelofibrosis or post polycythaemia p	Yes	TA386	23-Mar-16	21-Jun-16
RUX2	Ruxolitinib	For the treatment of polycythaemia vera for adult patients who are resistant to treatment with hydroxycarbamide or who cannot tolerate treatment with hydroxycarbamide where the following criteria have been met:	10. Rusoitlinib will otherwise be used as set out its Summary of Product Characteristics.  1. This applicants is being made by and the first cycle of systemic anti-cancer therapy with rusoitlinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a confirmed diagnosis of polycythaemia vera a Well-vision of the following criteria applying to this patient:  * age >60 years  * previous documented thrombosis (including transient ischaemic attack) or erythromelalgia or migraine (severe, recurrent, requiring medication and considered to be secondary to the PV) either after diagnosis of the PV or within the 10 years before diagnosis and regarded as being disease-related  * significant or symptomatic splenomegaly  * a platelet count exceeding 1000 x 1007 th, any point during the patient's disease  * diabetes or hypertension requiring pharmacological treatment for more than 6 months  4. The patient has been previously treated with hydroxycarbamide (HC) and is resistant to it or cannot tolerate treatment with it or is both resistant to it and intolerant of it.  **Note: the definitions of intolerance and resistance are those used by the European LeukaemiaNet (ELN) consensus.  **Please mark below which one of these scenarios applies to this patient:  ** the patient is sessiant to HC or  ** the patient is sessiant to HC or  ** the patient is been previously treated with rusoilinib or has received previous rusoilinib within the MAIC-PV trial or via a company compassionate access scheme and all the other criteria on this form are fulfilled.  ** The patient has not been previously treated with rusoilinib or has received previous rusoilinib within to many compassionate access scheme and all the other criteria on this from are fulfilled or  ** the patient has not been previously uncerted with rusoilinib or has received previous rusoilinib within the MAIC-PV trial and the benefit-risk ratio for continuing treatment remains positive and	Yes	TA921	18-Oct-23	16-Jan-24

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
SAC1_v1.1	Sacituzumab govitecan	For the treatment of patients with previously treated unresectable locally advanced or metastatic triple negative breast cancer where the following criteria have been met:	1. This application for sincturumb govitecan is been gmade by and the first cycle of systemic anti-cancer therapy with sacturumb govitecan will be prescribed by a consultant specialist specifically trained and accordited in the use of systemic therapy.  2. The patient has a histologically or optologically-confirmed diagnosis of breast cancer.  3. The patient has histologically or optologically-confirmed diagnosis of breast cancer.  4. The patient's breast cancer has had receptor analysis performed and this is negative for all of the following: the HER2 receptor, oestrogen receptor and progesterone receptor i.e. the patient has triple negative disease.  5. Either this patient has had 2 or more prior lines of systemic therapy specifically for the unresectable locally advanced or metastatic breast cancer indication and has also previously received adjuvant or necadipivant systemic therapy.  Please mark below which of these 2 clinical scenarios applies to this patient:  - this patient has been a control of these 2 clinical scenarios applies to this patient:  - this patient has only had 1 line of systemic therapy specifically for the unresectable locally advanced or metastatic breast cancer indication  - this patient has not been a controlled to the patient of the second of the patient has been treated with 1st line atecolizumab or pembrolizumab or the patient was technically eligible for 1st line atecolizumab or pembrolizumab or the patient was technically eligible for 1st line atecolizumab or pembrolizumab or pembrolizumab or the patient was technically eligible for 1st line atecolizumab or pembrolizumab or pembro	Yes	TA819	17-Aug-22	15-Nov-22

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
SELIN1	Selinexor in combination with bortezomib and dexamethasone	For the treatment of multiple myeloma in transplant ineligible patients who have had only 1 prior line of systemic therapy where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with sellneour in combination with bortezomib and deamerthasone will be prescribed by a consultant specialist specifically trained and according in the use of systemic and cancer therapy.  2. The patient has a diagnosis of multiple impelorus.  3. The prescribing discinsa understands that the combination of sellneour plus bortezomib and deamerthasone is notly for the specific 2nd line modified myelloms indication recommended by NCC.  Please tick hos below:  - this patient does not have a diagnosis of primary amyloidosis:  - this patient does not have a diagnosis of primary amyloidosis.  - this patient has received 3 and no more than 1 prior line of systemic treatment and that the numbering of a line of treatment is in accordance with the international Myelona Workshop Consensus recommendations for the uniform reporting of clinical trais (http://doi.org/10.1187/blood-2010-10.299467). A line of therapy is defined as one or more cycles of a planned treatment program. This may consist of one or more planned cycles of single-agent therapy or combination the large, as well as a sequence of treatments administrator of the international hyelena Workshop Consensus recommendations for the uniform response for them cell transplantation to proceed. A new line or demandation of the patient demandation is considered to be 1 line of the patient A new line or more planned reviews of a line of treatment program. This may consist of one or more planned cycles of single-agent therapy or combination therapy, as well as a sequence of treatments administrator of the planned manner (e) induction chemotherapy (between the patients when this observed by a need for additional treatment for the disease.  Please verify that the patient has only received 1 prior line of systemic therapy by ticking the box below:  - this patient has received 1 and no more than 1 prior line of systemic therapy by ticking the box below:  - this patient has received 1 and n	No	TA974	15-May-24	13-Aug-24

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
SELIN2	Selinexor in combination with dexamethasone	For the treatment of multiple myeloma in patients who have had at least 4 prior lines of systemic therapy and whose disease is refractory to at least 2 proteasome inhibitors, 2 immunomodulatory agents and an anti-CD38 monoclonal antibody and which has also demonstrated disease progression on the last therapy where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with selineary plus desamethssone will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The pattern has a diagnosis of multiple imploons.  3. The prescribed global consultance understance but the combination of selineary plus denamethssone is not funded for amyloidosis patients (with the exception of patients who have a proven diagnosis of progressive multiple myloidosis of amyloidosis and imploidosis) and that W16 funding for selineary plus decamendations is only for the specific 5th or more line multiple myloidosis and the combination of selineary plus decamendations of the myloidosis and the combination of selinear plus decamendations of the myloidosis.  **Repaired these not have a diagnosis of primary amyloidosis.**  **Inia patient does not have a diagnosis of primary amyloidosis.**  **Inia patient has received at least a fairer linear of progressive myloidosis and the combination of selinear plus decamendations of the myloidoma.**  **A. The patient has received at least a fairer linear of progressive myloidosis and the combination of selinear plus decamendations of the myloidoma.**  **A. The patient has received at least a fairer linear of systemic transport and the complex of single-agent the therapy of an ordinary of selections and the patients of the patients of selection and the patients of selection and the patients of selections are complex of selections.**  **A. The patient Systemic transport and the patients of selection and the patients of selections are complex of selections.**  **Pease state the number of lines of systemic therapy is considered by the patient by taking the appropriate box below.**  **Pease state the number of lines of systemic therapy is received by the patient by taking the appropriate box below.**  **Pease state the number of lines of systemic therapy will be patient by taking the appropriate box below.**  **Pease	- No	TA970	08-May-24	06-Aug-24

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
SELIN3	Selinexor in combination with bortezomib and dexamethasone	For the treatment of multiple myeloma in transplant ineligible patients who have had only 2 prior lines of systemic therapy and who are refractory to lenalidomide where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with sellensor in combination with bortecomb and decamenhasone will be prescribed by a consultant specialist specifically trained and according in the use of systemic anti-cancer therapy.  2. The patient has a diagnosis of multiple implementation of sellensor plus bortecomb and decamenhasone is not funded for anyloidosis patients (with the exception of patients who have a proven diagnosis of myeloma with a associated diagnosis of anyloidosis) and that Wish funding for sellensor plus bortecomb and decamenhasone is only for the specific 3rd line multiple myeloma indication recommended by MCC.  Plasar tick hos below:  - this patient does not have a diagnosis of primary anyloidosis.  - this patient does not have a diagnosis of primary anyloidosis.  - this patient does not have a diagnosis of primary anyloidosis.  - this patient has received 2 and no more than 2 prior lines of systemic treatment and that the numbering of a line of treatment is in accordance with the international Myeloma Workshop Consensus recommendations for the uniform apporting of clinical trais (https://doi.org/10.1182/bloco-2010-10.2998/17. A line of therapy is defined as one or more cycles of a planned treatment program. This may consist of one or more planned cycles of single-agent therapy or combination therapy, is well as a sequence of restmentation almost proceed. A new line of therapy also statis when a planned manner (le induction chemotherapy/chemotherapyis) when followed by stem cell transplantation is considered to be 1 line of therapy also statis when a planned manner (le induction chemotherapy/chemotherapyis) when followed by a need for additional treatment for the disease.  Please verify that the patient has only received 2 prior lines of systemic therapy by civiling the box below:  - this patient has received 2 and no more than 2 prior lines of systemic therapy by civiling the box below:  - this patient has received 2 and no more than 2 prior	No	TA974	15-May-24	13-Aug-24

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
SEL1	Selpercatinib		b. Ethner the patient has ontrerentated thyroid cancer (papiliary/foilicular/thurtic ceil) and has therefore deen treated with lenvatinilo or soratenio of the patient has anapiastic thyroid cancer in which case no previous IX i treatment requirement is necessary.  Please enter below as to the previous TXI therapy that the patient has received: - Inewatinib for differentiated thyroid cancer or - sorafenib for differentiated thyroid cancer or - has anaplastic thyroid cancer and hence no previous TXI therapy  - has anaplastic thyroid cancer and hence no previous TXI therapy  - The patient has an ECOG performance status (PS) of 0 or 1 or 2.  - Selpercatinib is being given as monotherapy.  - The patient has not previously received selpercatinib or any other TXI which targets the RET receptor unless the patient has received selpercatinib via a company early access scheme and the patient meets all the other criteria listed here.  - D. Selpercatinib is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment.  - The prescribing clinician is aware of the following issues as regards the administration of selpercatinib as detailed in its Summary of Product Characteristics (SPC):	- No	TA1038	12-Feb-25	started
			- the dosage of selpercatinib is according to body weight - selpercatinib has reduced solubility at a higher pH and hence precautions are necessary with the co-administration of proton pump inhibitors or H2 antagonists - selpercatinib has clinically important interactions with CYP3A inhibitors or CYP3A inducers  12. A formal medical review as to how selpercatinib is being tolerated and whether treatment with selpercatinib should continue or not will be scheduled to occur at least by the start of the third 4-weekly cycle of treatment.				
			13. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, a treatment break approval form will be completed to restart treatment.  14. Selpercatinib is to be otherwise used as set out in its Summary of Product Characteristics.				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
SEL2	Selpercatinib	For the treatment of adults or adolescents aged 12 years and older with previously treated RET mutant medulary thyroid cancer where the following criteria have been met:	2. This application is being made by and the first cycle of systemic anti-cancer therapy with selpercatinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient is an adult or an adolescent aged 12 years and older with a proven histological or cytological diagnosis of medullary thyroid cancer (there is a separate form SEL01 for selpercatinib in non-medullary thyroid cancer). Please enter below as to whether the patient is an adult or a dolescent aged 12 years or older.  1. The patient is an adult or  1. The patient is an adolescent aged 12 years or older  Note: if the patients is an adolescent aged 12 years or older  1. This patient's thyroid cancer has been documented as having a RET mutation as determined by a validated genomic test.  Please enter below as to which RET mutation is present in this patient's thyroid cancer:  1. This patient's thyroid cancer has been documented as having a RET mutation as determined by a validated genomic test.  1. The patient has been previously treated with cabozantinib or vanidatanib.  1. The patient has been previously treated with cabozantinib or vanidatanib.  1. The patient has been previously treated with cabozantinib or vanidatanib.  1. The patient has an ECOG performance status (PS) of 0 or 1 or 2.  1. Selipercatinib is being given as monotherapy.  1. The patient has not previously received selpercatinib or any other TRI which targets the RET receptor unless the patient has received selpercatinib via a company early access scheme and the patient meets all the other criteria listed here.  1. Selpercatinib is a coording to body weight  1. The patient has reduced solubility at a higher pH and hence precautions are necessary with the co-administration of proton pump inhibitors or H2 antagonists  1. Selpercatinib has reduced solubility at a higher pH and hence precautions or CYP3A inducers	No	TA1038	12-Feb-25	started
			10. A formal medical review as to how selpercatinib is being tolerated and whether treatment with selpercatinib should continue or not will be scheduled to occur at least by the start of the third 4-weekly cycle of treatment.  11. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, a treatment break approval form will be completed to restart treatment.  12. Selpercatinib is to be otherwise used as set out in its Summary of Product Characteristics.	-			

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
SEL3	Selpercatinib	Selpercatinib as monotherapy for the treatment of adult patients with advance non-small cell lung cancer (NSCLC) exhibiting a RET gene fusion and who have previously received immunotherapy and/or platinum-based chemotherapy where the following criteria have been met:	1. This application for seleptocration is being made by and the first cycle of systemic anti-cancer therapy with seleptocration will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has locally advanced or metastatic non-small cell lung cancer.  3. The patient has stategogacily or cyclogacily confirmed diagnosis of non-small cell lung cancer.  4. This patient's MSCL capites to this patient:  4. This patient's MSCL chaptes shown to hardour. 8. Eff game fusion as determined on a tumour tissue bioppy or a plasma specimen. (Isquid bioppy) or both.  4. This patient's MSCL chaptes shown to hardour. 8. Eff game fusion as determined on a tumour tissue bioppy or a plasma specimen (Isquid bioppy) or both.  4. This patient's MSCL chaptes shown to hardour. 8. Eff game fusion as determined on a tumour tissue bioppy or a plasma specimen (Isquid bioppy) or both.  4. This patient's MSCL chaptes shown to hardour. 8. Eff game fusion as determined to be in one of the categories as set out below:  4. In this patient's MSCL chaptes and plasma specimen.  5. This patient's MSCL between the patient's MSCL chaptes and plasma specimen.  5. This patient's MSCL between the patient's MSCL chaptes and plasma specimen.  6. This patient has proviously received immunotherapy and/or platinum-based chemotherapy for this locally advanced or metastatic MSCLC indication.  4. Provides and the patient has received is patient has received is patient has received size incombination returned of political metastatic machinerapy.  5. The patient has received size incombination returned of political metastatic MSCLC with or without 2 and ine cytotoxic chemotherapy or the patient has received size incombination returned of political metastatic MSCLC with or without 2 and ine cytotoxic chemotherapy or the patient has received size incombination or received size incombination or received size incombination and the patient machinerapy or locally advanced or metastatic MSCLC	No	TA1042	19-Feb-25	20-May-25

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
SELS	Selpercatinib		1. This application is being made by and the first cycle of systemic anti-cancer therapy with selpercatinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient has a proven histological or cytological diagnosis of non-medullary thyroid cancer (there is a separate form SEL6 for selpercatinib in medullary thyroid cancer previously untreated with any kinase inhibitor therapy).  Please enter below as to which type of thyroid cancer or  - anaplastic thyroid cancer or  - anaplastic thyroid cancer or  - anaplastic thyroid cancer or  - anaplastic thyroid cancer has been documented as having a RET fusion as determined by a validated genomic test.  Please enter below as to which is the RET fusion partner in this patient's thyroid cancer:  - CCDC6 or  - NCOA4 or  - another fusion partner  4. The patient is either an adult or an adolescent aged 12 years and older.  Please enter below as to which applies to this patient:  - the patient is an adult or	No	TA1039	12-Feb-25	started
		onoung citerio nave ocen mec	- the patient is an adolescent aged 12 years and older Note: if the patient is an adolescent, open growth plates should be monitored.  5. The patient is an adolescent, open growth plates should be monitored.  6. The patient is previously untreated with any kinase inhibitor unless the patient has received selpercatinib via a company early access scheme and the patient meets all the treatment criteria on this form.  7. The patient has an ECOG performance status (PS) of 0 or 1 or 2.  8. Selpercatinib is being given as monotherapy.  9. Selpercatinib is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment.  10. The prescribing clinician is aware of the following issues as regards the administration of selpercatinib as detailed in its Summary of Product Characteristics (SPC):  11. A formal medical review as to how selpercatinib is being tolerated and whether treatment with selpercatinib should continue or not will be scheduled to occur at least by the start of the third 4-weekly cycle of treatment.  12. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, the prescribing clinician will complete a treatment break approval form to restart treatment.  13. Selpercatinib is to be otherwise used as set out in its Summary of Product Characteristics.				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
SEL6	Selpercatinib	For the treatment of adults or adolescents aged 12 years and older with RET mutant medullary thyroid cancer previously UNTREATED with any kinase inhibitor therapy where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with selpercatinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient is an adult or an adolescent aged 12 years and older with a proven histological or cytological diagnosis of medullary thyroid cancer (there is a separate form SEL5 for selpercatinib in non-medullary thyroid cancer previously untreated with any kinase inhibitor therapy).  Please enter below as to which applies to this patient: - the patient is an adolescent aged 12 years and older Note: if the patient is an adolescent aged 12 years and older Note: if the patient is an adolescent aged 12 years and older Note: if the patient is an adolescent aged 12 years and older Note: if the patient is an adolescent, open growth plates should be monitored.  3. This patient's thyroid cancer has been documented as having a RET mutation as determined by a validated genomic test.  Please enter below as to which RET mutation is present in this patient's thyroid cancer:  - mostar mutation or - another mutation or - v804M/L mutation or - another mutation  4. The patient is previously untreated with any kinase inhibitor unless the patient has received selpercatinib via a company early access scheme and the patient meets all the treatment criteria on this form.  5. The patient has an ECOG performance status (PS) of 0 or 1 or 2.  6. Selpercatinib is being given as monotherapy.  7. Selpercatinib is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment.  8. The prescribing clinician is aware of the following issues as regards the administration of selpercatinib as detailed in its Summary of Product Characteristics (SPC): - the dosage of selpercatinib is according to body weight - selpercatinib has reduced solubility at a higher pH and hence precautions are necessary with the co-administration of proton pump inhibitors or H2 antagonists - selperc	No	TA1039	12-Feb-25	13-May-25

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
SOR2	Sorafenib	The treatment of differentiated thyroid cancer after radioactive iodine where the following criteria are met:	1. This application is made by and the first cycle of systemic anti-cancer therapy with sorafenib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. This patient has a confirmed histological diagnosis of differentiated thyroid carcinoma (papillary or follicular or Hurtle cell type)  3. The patient has either metastatic disease or inoperable locally advanced disease  4. The disease is refractory to radioactive iodine  5. The disease is progressive and is either symptomatic or imminently likely to become symptomatic  6. The patient is treatment naïve to both lenvatinib and sorafenib unless the patient has had to discontinue lenvatinib within 3 months of starting lenvatinib because of toxicity (ie there is lenvatinib toxicity which cannot be managed by dose delay or dose modification) and there has been no disease progression whilst on lenvatinib.  Note: Sequential use of sorafenib and then lenvatinib is only funded if the patient has to discontinue sorafenib because of intolerance within 3 months of its start and if the disease has not progressed whilst the patient is on sorafenib. The use of sorafenib after disease progression on or after lenvatinib is not funded and vice versa.  7. The patient has an ECOS performance status of 0 or 1 or 2.  8. Sorafenib is to be continued as long as clinical benefit is observed or until there is unacceptable toxicity or patient choice to stop treatment.  9. A formal medical review as to whether treatment with sorafenib should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment  10. No treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve)  11. Sorafenib is to be otherwise used as set out in its Summary of Product Characteristics	Yes	TA535	08-Aug-18	06-Nov-18
SOR3	Sorafenib monotherapy	Treatment of Child-Pugh A locally advanced or metastatic hepatocellular carcinoma where the following criteria are met:	1. An application has been made by and the first cycle of systemic anti-cancer therapy is to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. ONE of the following applies to the patient: the patient has a confirmed histological diagnosis of hepatocellular carcinoma or a biopsy is deemed to be very high risk or technically not feasible in the patient AND the criteria below are met:  a. The decision not to biopsy has been made and documented by a specialist HCC MDM  b. The tumour meets the non-invasive diagnostic criteria of hepatocellular carcinoma*  c. Data is submitted as part of the ongoing Sorafenib Audit 2.  It is expected that OPTION 2 will only apply in exceptional circumstances and it should be noted that responses will be reviewed regularly to ensure that this is the case.  *EASE-FORTC Clinical Practice Guidelines: Management, Journal of Hepatology 2012 vol. 56 p 908–943. Non-invasive criteria can only be applied to circhotic patients and are based on imaging techniques obtained by 4-phase multidetector CT scan or dynamic contrast-enhanced MRI. Diagnosis should be based on the identification of the typical hallmark of HCC (hypervascular in the arterial phase with washout in the portal venous or delayed phases). While one imaging technique is required for nodules beyond 1 cm in diameter a more conservative approach with 2 techniques is recommended in suboptimal settings.  3. Patient must have either metastatic disease or locally advanced disease that is ineligible for or failed surgical or locoregional therapies  4. Either:  - the patient has not received any previous systemic therapy for hepatocellular carcinoma (option 1) or  - the patient has had to discontinue lenvatinib within 3 months of starting lenvatinib and solely because of toxicity (i.e. there was lenvatinib toxicity which could not be managed by dose delay or dose modification) and there has been not disease progression whilst no lenvatinib (potion 2) or  - if the pat	Yes	TA474	06-Sep-17	05-Dec-17

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
SOR5	Sorafenib	ITD) acute myeloid leukaemia (AML) post allogeneic haematopoietic stem cell	1. This application is being made by and the first cycle of systemic anti-cancer therapy with sorafenib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a diagnosis of FLT3-internal Tandem Duplication (FLT3-ITD) mutation acute myeloid leukaemia (AML).  3. The patient is aged 18 and over.  4. Sorafenib is not licensed for FLT3-ITD mutation AML maintenance therapy post allogeneic haematopoietic stem cell transplantation (allo-HSCT) and therefore Trust policy regarding unlicensed medicines has been followed.  5. The patient has been discussed by a relevant specialist MDT and it has been agreed that sorafenib is the most appropriate therapy.  6. Sorafenib in this indication is to be used as monotherapy and not combined with any other maintenance therapy and that the patient will receive the recommended dosing of sorafenib as outlined in the NH5 England Clinical Commissioning Policy and the product Summary of Product Characteristics.  7. The patient meets all of the following eligibility criteria:  o has undergone allogeneic haematopoietic stem cell transplantation AND  o Exhibits adequate engraftment (absolute neutrophil count of at least 10 x 10°/L and a non-transfused platelet count of at least 30 x 10°/L) at the time of sorafenib initiation.  8. The patient one of the following eligibility criteria:  o Individuals with contraindications to sorafenib, as outlined in the summary of product characteristics (SPC) OR  o Uncontrolled graft versus host disease (SWH) OR  o Persistent liver dyfunction (total billirubin twice or more the upper limit of normal [ULN] or alanine aminotransferase or aspartate aminotransferase twice or more the ULN) OR  o Persistent rever dyfunction (total billirubin twice or more the upper limit of normal [ULN] or alanine aminotransferase or aspartate aminotransferase twice or more the ULN OR or Persistent rever dyfunction (total billirubin twice or more the upper limit of normal [ULN] or alanine	No	NHSE Policy: URN2262	N/A	06-Nov-23
SOR6	Sorafenib	Sorafenib maintenance for the treatment of FLT3-Internal Tandem Duplication (FLT3 ITD) acute myeloid leukaemia (AML) post allogeneic haematopoietic stem cell transplantation (allo-HSCT) IN POST-PUBESCENT CHILDREN where the following criteria are met:	13. Sorafenib will otherwise be used as set out in its Summary of Product Characteristics (SPC).  1. An application has being made by and the first cycle of systemic anti-cancer therapy with sorafenib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient is a post-pubescent child receiving access under the Medicines for Children policy.  4. Sorafenib is not licensed for FLT3-ITD mutation ANU maintenance therapy post allogeneic heamatopoletic stem cell transplantation (allo-HSCT) and therefore Trust policy regarding unlicensed medicines has been followed.  5. The patient has been discussed by a relevant specialist MDT and it has been agreed that sorafenib is the most appropriate therapy. This MDT must include at least two consultants with experience in the treatment of FLT3-ITD ANL of whom at least one must be a consultant paediatrician. The MDT should also include a paediatric pharmacist and other professional groups appropriate to the disease area.  6. Looffirm that sorafenib in this indication is to be used as monotherapy and not combined with any other maintenance therapy and that the patient will receive the recommended dosing of sorafenib as outlined in the NHS England Clinical Commissioning Policy and the product's Summary of Product Characteristics.  7. The patient meets all of the following eligibility criteria:  o has undergone allogeneic haematopoletic stem cell transplantation AND  o Exhibits adequate engraftment (absolute neutrophil count of at least 1.0 x 10°/L and a non-transfused platelet count of at least 3.0 x 10°/L) at the time of sorafenib initiation.  8. The patient meets all of the following eligibility criteria:  o Individuals with contraindications to sorafenib, as outlined in the summary of product characteristics (SPC) OR  o Uncontrolled graft versus host disease (GMTD) OR  o Persistent renal dysfunction (total billimibin twice or of the following events and produce of the following events of the fol	No	NHSE Policy: URN2262	N/A	06-Dec-23

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
SUN1	Sunitinib	The treatment of unresectable or metastatic neuroendocrine tumours of pancreatic origin with disease progression where all the following criteria are met:	1. An application has been made by and the first cycle of systemic anti-cancer therapy is to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The patient has histopathologically proven well differentiated neuroendocrine tumour of pancreatic origin  3. The patient has exhibited disease progression in past 12 months  5. The patient has a performance status of 0-1  6. The patient has a performance status of 0-1  6. The patient has had no previous treatment with a tyrosine kinase inhibitor.  7. No planned treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve).*  *Requests for continuation of treatment after unplanned treatment breaks over this duration should be made via the treatment break approval process  8. Sunitinib will otherwise be used as set out in its Summary of Product Characteristics (SPC).	Yes	TA449	13-May-17	26-Sep-17

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
TAL1	Talazoparib monotherapy	Talazoparib as monotherapy for treatmen of adults with deleterious or suspected deleterious germline BRA1 or 2 mutations who have HER-2 negative locally advanced or metastatic breast cancer previously treated with an anthracycline and/or taxane in the adjuvant/neoadjuvant/advanced disease settings and also treated with prior endocrine-based therapy if the patient ha hormone-receptor positive disease where the following criteria have been met:	1. This application is both being made by and the first cycle of systemic anti-cancer therapy with talasoparith monotherapy will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. This patient has a proven histological diagnosis of MRR 2 negative breast cancer.  Note: Instance and the control of the contro	No	TA952	21-Feb-24	21-May-24

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
TAU1	Talimogene Laherparepvec	Talimogene laherparepvec for treating unresectable metastatic melanoma	1. It confirm that an application has been made and the first treatment will be prescribed and administered by a consultant specialist experienced in the treatment of melanoma  2. I confirm this treatment will be given by a specialist trained to give intra-lesional injections of talimogene.  3. I confirm the patient has cutaneous, subcutaneous or notal deposit(s) of melanoma which is/are suitable for direct injection but is/are not surgically resectable.  4. I confirm the patient has stage lilb, stage lic or stage IVM1a disease according to the AICC stage criteria of 2009 7th edition and if stage IVM1a disease (in emtastases to the skin, subcutaneous tissues or distant lymph nodes) has a normal serum LDH.  5. I confirm the patient has no bone, brain, lung or any other visceral secondaries and if stage IVM1a disease, the serum LDH is not elevated.  6. I confirm talimogene has been sanctioned by a specialist melanoma multidisciplinary team which includes an oncologist and a surgeon with expertise in the management of metastatic and locally advanced melanoma, respectively.  7. I confirm that talimogene part appropriate for this patient as systemically administered minuncherapies or approved targeted therapies are not considered the best option by the specialist melanoma multidisciplinary team meeting which includes an oncologist and a surgeon with expertise in the management of metastatic and locally advanced melanoma, respectively.  8. I confirm that talimogene will only be administered as a single agent and not in combination with systemic therapies eg chemotherapy, targeted agents or immunotherapy unless this is within the context of a Health Research Authority clinical trial.  9. I confirm the patient will receive the licensed dose and frequency of talimogene laherparepovec	No	TA410	28-Sep-16	28-Dec-16
Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application for tebentafusp monotherapy is both being made by and the first cycle of systemic anti-cancer therapy with tebentafusp will be prescribed by a consultant melanoma specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient is an adult with a histologically proven diagnosis of uveal melanoma.  3. The patient has unrescrable or metastatic uveal melanoma.  5. The patient has urnescrable or metastatic uveal melanoma.  5. The patient does not have symptomatic or untreated brain metastases.  6. The patient has either been previously treated with any prior systemic therapy or not including if the patient has received tebentafusp via a company early access scheme and all other treatment criteria on this form apply.  Please mark below which clinical scenario applies to this patient:  - the patient has been treated with any prior systemic therapy or tebentafusp  - the patient has been treated with prior checkpoint inhibitor systemic therapy and has not received prior tebentafusp  - the patient has been treated with prior checkpoint inhibitor systemic therapy and has not received prior tebentafusp and all other treatment criteria on this form apply				
TEB1	Tebentafusp	Tebentafusp as monotherapy for adult patients with human leukocyte antigen HLA-470:201 positive unrescrable or metastatic uveal melanoma where the following criteria have been met:	9. The treating hospital has facilities (including those for resuscitation) to manage severe reactions to tebentafusp including cytokine release syndrome (CRS).  10. The prescribing clinician and the treating team are aware of the risks and grading of cytokine release syndrome (CRS),  Its monitoring and management as illustrated in Table 1 of section 4.2 of the tebentafusp Summary of Product Characteristics and both I and the treating team have all undergone training in these clinical issues.  11. Clear arrangements have been made for the patient to be monitored as an inpatient for signs and symptoms of toxicities including CRS for 16 hours after administration of the first 3 x weekly doses of tebentafusp.  12. The prescribing clinician and the treating team are aware that if any grade 3 or 4 hypotension occurs during any of the first 3 infusions, the patient will be monitored every hour for the next 4 hours in an outpatient setting for the 13. There is immediate access to treatment with tocilizumab if required to manage CRS.  14. The patient will be treated with tebentafusp until there is clear evidence of progressive disease or the occurrence of excessive toxicity or the withdrawal of patient consent, whichever is the sooner.	No	TA1027	09-Jan-25	09-Apr-25
			15. A formal medical review as to how tebentafusp is being tolerated and whether treatment with tebentafusp should continue or not will be scheduled to occur at least by the end of the first 4 weeks of treatment.  16. When a treatment break of more than 6 weeks beyond the expected weekly cycle length is needed, a treatment break approval form will be completed to restart treatment.  17. Tebentafusp will be otherwise used as set out in its Summary of Product Characteristics (SPC).				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
TEC1	Teclistamab	For the treatment of relapsed or refractory myeloma in adult patients who have relapsed or are refractory to their last anti-myeloma regimen AND have received at least 3 prior lines of systemic therapies which must have included at least one proteasome inhibitor, at least one immune-modulatory agent and at least one anti-CD38 antibody and where the following criteria have been met:	1. This application for tecilization binomorphisms just both being made by and the first cycle of systemic and iscancer therapy with techtiams will be prescribed by a consultant specialist specifically trained and accordinate in the use of systems and studies with prevent diagnosis of multiple implement.  Volume patients will be the relevant be bothow.  Please tick the relevant be bothow.  It is gained does not law a diagnosis of primary amyloidosis or the training of the studies o	No	TA1015	13-Nov-24	11-Feb-25

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Blueteq Form ref	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
TEC1	Teclistamab	For the treatment of relapsed or refractory myeloma in adult patients who have relapsed or are refractory to their last anti-myeloma regimen AND have received at least 3 prior lines of systemic therapies which must have included at least one protessome inhibitor, at least one inmune-modulatory agent and dat least one anti-CDB antibody and where the following criteria have been met:	11. The patient has been treated with a 8CMA-targeted antibody drug conjugate (such as belantamab mafodotin).  Please confirm which situation applies to this patient:		TA1015	13-Nov-24	11-Feb-25

v1.379 25-Nov-2025

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
TEP1	Tepotinib	Tepotinib as monotherapy for the treatment of adult patients with untreated advanced/metastatic non-small cell lung cancer (MSCLC) harbouring mesenchymla-eiphtleial transition (MET) exon 14 skipping alterations where the following criteria are met:	1. This application for tepotinib is being made by and the first cycle of systemic anti-cancer therapy with tepotinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has locally advanced or metastatic non-small cell lung cancer.  Please indicate below whether the patient has non-squamous or squamous NSCLC:  - non-squamous NSCLC or  - squamous NSCLC  3. The patient has histological or cytological evidence of NSCLC that carries a MET exon 14 skipping alteration based on a validated test OR there is documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC AND there is an informative circulating free DNA test result confirming the presence of a MET exon 14 skipping alteration.  Please mark below on which basis the diagnosis of a MET exon 14 skipping alteration positive NSCLC has been made in this patient:  - Documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC and there is an informative circulating free DNA test result confirming the presence of a MET exon 14 skipping alteration  4. The patient's lung cancer is of EGFR wild type and is also negative for both ALK and ROS1 gene rearrangements.  5. This patient is treatment-naive as regards to systemic therapy for the locally advanced or metastatic NSCLC indication.  6. The patient has not been previously treated with a drug specifically targeting a MET exon 14 skipping alteration unless the patient received tepotinib via the EAMS program and the patient meets all the other treatment criteria on this form.  7. The patient has no theorem previously treated with a drug specifically targeting a MET exon 14 skipping alteration unless the patient received tepotinib via the EAMS program and the patient meets all the other treatment criteria on this form.  7. The patient either has no known brain meetastases or if the patient does have br	No No	TA789	18-May-22	started  17-Jun-22
			scheduled to occur at least by the end of the second month of therapy.  14. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment, including as appropriate if the patient has had an extended break on account of Covid-19.  15. Tepotinib will otherwise be used as set out in its Summary of Product Characteristics (SPC).	ie			

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application for tepotinib is being made by and the first cycle of systemic anti-cancer therapy with tepotinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has locally advanced or metastatic non-small cell lung cancer.  Please indicate below whether the patient has non-squamous or squamous NSCLC:  - non-squamous NSCLC or  - squamous NSCLC  3. The patient has histological or cytological evidence of NSCLC that carries a MET exon 14 skipping alteration based on a validated test OR there is documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC AND there is an informative circulating free DNA test result confirming the presence of a MET exon 14 skipping alteration.  Please mark below on which basis the diagnosis of a MET exon 14 skipping alteration NSCLC has been made in this patient:  - Histological or cytological evidence.  - Documented agreement by the lung MDT that the radiological appearances are in keeping with locally advanced or metastatic NSCLC and there is an informative circulating free DNA test result confirming the presence of a MET exon 14 skipping alteration.				
TEP2	Tepotinib	Tepotinib as monotherapy for the treatment of adult patients with previously treated advanced/metastatic non-small cell lung cancer (NSCLC) harbouring mesenchymal-epithelial transition (MET) exon 14 skipping alterations where the following criteria	4. The patient's lung cancer is of EGFR wild type and is also negative for both ALK and ROS1 gene rearrangements.  5. This patient has previously received systemic therapy for the locally advanced or metastatic NSCLC indication: As regards the previous treatment received by the patient, please mark which of these 5 scenarios below applies to this patient: - the only treatment that the patient has received is platinum-based cytotoxic chemotherapy for locally advanced or metastatic NSCLC with or without 2nd line cytotoxic chemotherapy or - the only treatment that the patient has received is 1st line immunotherapy monotherapy for the locally advanced or metastatic NSCLC indication or - the patient has received the 1st line combination treatment of platinum doublet chemotherapy plus immunotherapy for the locally advanced or metastatic NSCLC indication with or without 2nd line cytotoxic chemotherapy or - the patient has received 1st line immunotherapy monotherapy for the locally advanced or metastatic NSCLC indication followed by 2nd line cytotoxic chemotherapy or - the patient has received 1st line immunotherapy for locally advanced or metastatic NSCLC indication followed by 2nd line cytotoxic chemotherapy or - the patient has received 1st line platinum-based cytotoxic chemotherapy for locally advanced or metastatic NSCLC followed by 2nd line immunotherapy with or without further cytotoxic chemotherapy  6. The patient has not been previously treated with a drug specifically targeting a MET exon 14 skipping alteration unless the patient received tepotinib via the EAMS program and the patient meets all the other treatment criteria on	No	TA789	18-May-22	17-Jun-22
		are met:	7. The patient has not been previously deated with a drug specifically targeting a MET exon 14 supping alteration unless the patient received tepotinio via the Early Signature and the patient meets all the other treatment Criteria on this form.  7. The patient has an ECOG performance status (PS) score of 0 or 1.  8. The patient has no known brain metastases or if the patient does have brain metastases then the patient is symptomatically stable before staring tepotinib.  Please mark below the status with respect to known brain/CNS metastases:  - the patient has never had known brain/CNS metastases  - the patient has had brain/CNS metastases treated before with surgery/radiotherapy and is currently symptomatically stable  - the patient has brain secondaries which have not been treated with surgery/radiotherapy and is currently symptomatically stable				
			9. Tepotinib will be used as monotherapy. 10. The prescribing clinician is aware of the side-effects of tepotinib including the risk of developing oedema, interstitial lung disease and hepatotoxicity. 11. The prescribing clinician is aware of the potential drug interactions that may occur with tepotinib as set out in tepotinib's Summary of Product Characteristics (SPC) including the potential interaction with metformin. 12. The patient will be treated until loss of clinical benefit or excessive toxicity or patient choice to discontinue treatment whichever is the sooner. 13. A formal medical review as to how tepotinib is being tolerated will be done before the start of the second month of treatment and the next review to determine whether treatment with tepotinib should continue or not will be scheduled to occur at least by the end of the second month of therapy. 14. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment, including as appropriate if the patient has had an extended break on account of Covid-19.				

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lueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
TISO1a	Tisageniecieucel	will relates to the subsequent intusion of CAR-T cells and this will be available after submission of the first part. The second part of the form (TiSO1a) can only be completed as a continuation of this first part of the form (TiSO1a) and must be completed on infusion of CAR-T cells otherwise the treating Trust will not be	1. This application is being made by and that leurapheresis for and treatment with tisagenicefuscien-modified CART cells will be initiated by a consultant haematologist specifically trained and accretified in the use of systemic anticeacher therapy and voring in an accredite CART cell multidisciplinary teams.  2. The patient has required to refractory as lineage acute lymphobbastic leukaemia and CART Cell multidisciplinary teams.  2. The patient has required for refractory Bineage acute lymphobbastic leukaemia (ALI).  Please tick appropriate box as to which type of ALL the patient has:  Publisdisphia chromosome positive ALI  3. The patient fulfils one of the following clinical accessing relating to the definition of relapsed or refractory ALI:  2. And or more bown marrow relapse following conventional dises of chemotherapy/immocidural antibody therapy. On  2. And or more bown marrow relapse following conventional dises of chemotherapy/immocidural antibody therapy. On  2. For immary refractory disease in on a children's accomplete remission after 2 cycles of standard chemotherapy for newly diagnosed ALI. On  2. For immary refractory disease in on a children's accomplete remission after 2 cycles of standard chemotherapy for newly diagnosed ALI. On  2. For immary refractory disease in on a childrenge a complete remission after 2 cycles of standard chemotherapy for newly diagnosed ALI. On  3. The patient has Philadelphia positive ALI that is refractory to primary chemotherapy or has relapsed post transplant or is in 2nd or greateer relapse despite treatment with standard chemotherapy plus TII therapy OR  4. Having fulfilled and ticked one of the criteria in box 3 above, the patient at the time of demonstration of such refractory/relapsed disease and thus consideration for potential treatment with tisageniceducel either had a bone  marrow with both flow cytometry detectable ALI and CD19 ALI positivity in the bone marrow or in the case of an isolated CNS relapse had both flow cytometry detectable ALI and CD19 ALI pos	Yes	TA975	15-May-24	13-Aug-24
TISO1b	Tisagenlecleucel	under where the following criteria are met:  Note: This second part of the form is to document the date of infusion of CAR-T cell therapy and for registration of this infusion with NHS Englands os that the treating Trust is reimbursed for the cost of tisagentedeucel. Three is a first part of the form for the approval of leucapheresis and manufacture of CAR-T cells which has already been completed (TISO1a). This	listed here.  1. This application for continuation is being made by and treatment with tisagenlecleucel-modified CAR T cells will be initiated by a consultant haematologist specifically trained and accredited in the use of systemic anti-cancer therapy and working in an accredited CAR T cell treatment centre and who is a member of the National CAR T Clinical Panel for acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's acute lymphoblastic leukaemia and a member of the treating Trust's a	Yes	TA975	15-May-24	13-Aug-24

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
TIV1	Tîvozanib	The treatment of advanced renal cell carcinoma where all the following criteria are met:	1. This patient has a bistologically- or cytologically-proven diagnosis of renal cell carcinoma (RCC) which either has a clear cell component or is one of the types of RCC as indicated below.  Please indicate below which RCC histology applies to this patient:  - RCC with a clear cell component or is one of the types of RCC as indicated below.  Please indicate below which RCC histology applies to this patient:  - ArcC with a clear cell component or is one of the types of RCC as indicated below.  - RCC with a clear cell component or is one of the types of RCC as indicated below.  - ArcC with a clear cell component or is one of the types of RCC as indicated below.  - ArcC with a clear cell component or is one of the types of RCC as indicated below.  - ArcC with a clear cell component or is one of the types of RCC as indicated below.  - ArcC with a clear cell component or is one of the types of RCC as indicated below.  - ArcC with a clear the RCC get a clear collecting duct RCC or - medium place (RCC or - medium) and place (RCC or - medium) and place (RCC or - multipolitic volte RCC  No	TAS12	21-Mar-18	19-Jun-18	

lueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application is made by and the first cycle of systemic anti-cancer therapy with trametinib in combination with dabrafenib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy	-			
			2. This patient has a confirmed histological diagnosis of malignant melanoma which is BRAF V600 mutation positive				
			3. The patient has unresectable stage III or stage IV disease that has been staged according to the AICC 8th edition	1			
TRADAB1	Trametinib and	Trametinib in combination with dabrafenib for treating unresectable or	4. The patient is treatment naïve to BRAF V600 and MEK inhibitors for malignant melanoma unless either the patient has previously received adjuvant dabrafenib and trametinib and did not progress during such therapy or has received a sufficient trial of encorafenib plus binimetinib for advanced disease and this has had to be stopped solely as a consequence of persistent dose-limiting toxicity and in the documented absence of disease progression.  Note: sequential treatment is not commissioned with encorafenib plus binimetinib and then on disease progression with dabrafenib plus trametinib.	No	TA396	22-Jun-16	20-Sep-16
TRADABI	Dabrafenib	metastatic melanoma where the following	5. The patient has sufficient ECOG performance status to tolerate treatment with the combination of trametinib plus dabrafenib	NO	1A390	22-Jun-16	20-3ep-10
		criteria have been met:	6. Treatment with trametinib in combination with dabrafenib will be continued until loss of clinical benefit or unacceptable toxicity or withdrawal of patient consent. The only exception to this is for patients enrolled in the NIHR-approved INTERIM trial in which intermittent treatment is allowed and can be given in the experimental arm				
			7. A formal medical review as to whether treatment with trametinib in combination with dabrafenib should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment				
			8. No treatment breaks of more than 6 weeks beyond the expected 4-weekly cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve)*  *Requests for continuation of treatment after unplanned treatment breaks over this duration should be made via the treatment break approval process.				
			9. Trametinib in combination with dabrafenib is to be otherwise used as set out in their respective Summaries of Product Characteristics				
			1. This application is made by and the first cycle of systemic anti-cancer therapy with dabrafenib in combination with trametinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy	-			
			2. This patient has a confirmed histological diagnosis of malignant melanoma which is BRAF V600 mutation positive				
			3. The patient has disease that has been staged as stage III disease according to the AICC 8th edition				
			4. This stage III disease has been completely resected either via sentinel lymph node biopsy ('sentinel lymphadenectomy') or when indicated via completion lymph node dissection and/or there has been complete resection of intransit metastases.				
			5. The patient is treatment naïve to systemic therapy for malignant melanoma and in particular has not previously received any BRAF V600 inhibitors or MEK inhibitors or immunotherapy with any check point inhibitors				
TRADAB2	Trametinib and Dabrafenib in combination with trametinib G. The prescribing clinician has discussed with the patient the benefits and toxicities of adjuvant trametinib and dabrafenib in stage III disease, the said 10 year melanoma-specific survival probabilities with routine surveillance are 93% and 88%, respectively respectively criteria are met:    Dabrafenib in combination with trametinib G. The prescribing clinician has discussed with the patient the benefits and toxicities of adjuvant trametinib and dabrafenib in stage III disease, the said 10 year melanoma-specific survival probabilities with routine surveillance are 93% and 88%, respectively for stage III disease, the 5 and 10 year figures are 83% and 77%, respectively    For stage III disease, the 5 and 10 year figures are 69% and 60%, respectively	- for stage IIIA disease, the 5 and 10 year melanoma-specific survival probabilities with routine surveillance are 93% and 88%, respectively - for stage IIIB disease, the 5 and 10 year figures are 83% and 77%, respectively - for stage IIIC disease, the 5 and 10 year figures are 69% and 60%, respectively	No	TA544	17-Oct-18	15-Jan-19	
			- for stage IIID disease, the 5 and 10 year figures are 32% and 24%, respectively.				
			7. The patient has an ECOG performance status of either 0 or 1				
			8. Treatment with dabrafenib in combination with trametinib will be continued for a maximum of 12 months from the start of treatment in the absence of disease recurrence, unacceptable toxicity or withdrawal of patient consent				
			9. A formal medical review as to whether treatment with dabrafenib in combination with trametinib should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment				
			10. No treatment breaks of more than 6 weeks beyond the expected 4-weekly cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve)*  *Requests for continuation of treatment after unplanned treatment breaks over this duration should be made via the treatment break approval process.				
			11. Dabrafenib in combination with trametinib is to be otherwise used as set out in their respective Summaries of Product Characteristics.				
			1. This application for dabrafenib and trametinib for BRAF V600-mutated anaplastic thyroid cancer (ATC) is being made by and the first cycle of dabrafenib and trametinib for BRAF V600-mutated ATC will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anticancer therapy.				
			2. The patient has been diagnosed with locally advanced inoperable anaplastic thyroid cancer.				
	Trametinib and	Dabrafenib in combination with trametinib for BRAF V600-mutated anaplastic thyroid	3. The patient has been tested for and has a confirmed BRAF V600 mutation.		NHSE Policy:		
TRADAB3	Dabrafenib	cancer (ATC) for ADULT patients where	4. The patient has a performance status of 0 or 1 or 2.	No	221006P	N/A	21-Oct-22
		the following criteria have been met:	5. Dabrafenib and trametinib for BRAF V600-mutated anaplastic thyroid cancer are to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment.	-			
			6. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, a treatment break approval form will be completed to restart treatment.  Debuffeible date treatment by the property of the	-			
		7. Dabra	7. Dabrafenib and trametinib will be otherwise used as set out in their respective Summary of Product Characteristics (SPCs).  8. Trust policy regarding the use of unlicensed (off-label) treatments has been followed as these drugs in this treatment are not licensed in this indication.	-			

lueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
			1. This application for trastuzumab emtansine as adjuvant chemotherapy is being made by and the first cycle of adjuvant trastuzumab emtansine will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has histologically documented breast cancer which is HER2 3+ by immunohistochemistry and/or has a ratio of ≥2.0 by in situ hybridisation.				
			3. The patient has been diagnosed with early breast cancer and this has been adequately excised.				
			4. Prior to neoadjuvant chemotherapy the patient had clinical stage T1-T4, nodal stage N0-3 and metastasis stage M0 disease.				
			5. The patient has been previously treated with at least 16 weeks of neoadjuvant cytotoxic chemotherapy which incorporated a minimum of at least 9 weeks of taxane-based chemotherapy and 9 weeks of HER2-targeted therapy unless entered into the ROSCO trial or was considered potentially eligible for the HER2 RADICAL trial.  Please tick below which option applies:  - At least 16 weeks of neoadjuvant cytotoxic chemotherapy which incorporated a minimum of at least 9 weeks of taxane-based chemotherapy and at least 9 weeks of HER2-targeted therapy or  - The patient was enrolled into the ROSCO trial (UKCRN Study ID19069) and was treated with 4 cycles of neoadjuvant chemotherapy plus trastuzumab with or without pertuzumab but did not achieve a pathological complete response				
			and has therefore received 4 cycles of adjuvant chemotherapy with trastuzumab with or without pertuzumab or - The patient was potentially legible for the HERS ADIACL trial (LKCRN Study In 13182) and was retarted with at least 12 weeks of taxane-based chemotherapy with trastuzumab and pertuzumab but did not achieve a pathological complete response and has therefore received at least 9 weeks of anthracycline-based adjuvant treatment	No TA632			
TRA2	Trastuzumab emtansine	As adjuvant therapy for patients with HER2-positive early breast cancer who have residual invasive disease following the combination of taxane-based and HER2-targeted neoadjuvant systemic therapy and surgery where the following	6. The patient has documented residual disease after neoadjuvant chemotherapy and HER2-directed treatment and that one of the following scenarios applies to this patient as to the documented residual invasive disease after completion of neoadjuvant therapy and surgery:  - the patient had residual invasive disease in the breast only or  - the patient had residual invasive disease in the lymph nodes only or  - the patient had residual invasive disease in both the breast and lymph nodes.  Note: trastuzumab emtansine as adjuvant treatment is only NICE-recommended and NHS England-commissioned in patients with documented residual disease invasive disease after completion of neoadjuvant chemotherapy and surgery.		TA632	10-Jun-20	08-Sep-20
		criteria have been met:	2. Adjuvant trastuzumab emtansine will be used as monotherapy.  8. Trastuzumab emtansine is the only HER2-directed therapy to be given after surgery i.e. no adjuvant trastuzumab/pertuzumab has been administered since surgery with the exception of patients enrolled in the ROSCO clinical trial. It is acknowledged that post-surgery patients may have received one cycle of adjuvant pertuzumab and trastuzumab whilst awaiting the pathology results to confirm the status of axillary lymph node involvement and any residual				
			disease  9. A maximum of 14 cycles of trastuzumab emtansine will be administered as adjuvant therapy unless there is evidence of progressive disease or unacceptable toxicity or withdrawal of patient consent.  If trastuzumab emtansine has to be discontinued early, and without disease progression, completion of the intended adjuvant reatment duration up to 14 cycles of adjuvant HER2-directed therapy can be done with trastuzumab (if lymph node negative) or trastuzumab plus pertuzumab (if lymph node positive).  Note: A maximum of 18 cycles of HER2-directed therapy (neadplyinant plus adjuvant) are funded provided all other criteria are met.  10. The patient has an ECOG performance status of 0 or 1.  11. The left ventricular ejection fraction prior to commencing adjuvant treatment with trastuzumab emtansine remains ≥50%.				
			13. Trastuzumab emtansine will be otherwise used as set out in its Summary of Product Characteristics (SPC).				
			1. An application has been made by and the first cycle of systemic anti-cancer therapy is to be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy				
			2. Progression of her-2 positive locally advanced or metastatic breast cancer 3. Progression during or after the most recent treatment for advanced stage disease or within 6 months of completing treatment for early stage disease 4. Previous treatment with a taxane OR capecitabine.				
		The treatment of HER2-positive locally	4. Previous treatment with rastructumab  5. Previous treatment with trastructumab  5. Previous treatment with trastructumab				
TRA1	Trastuzumab Emtansine	advanced/ unresectable or metastatic	6. Perfomance statau of 0, 1 or 2	Yes	TA458	19-Jul-17	17-Oct-17
	Trustazamus Emtansme	(Stage IV) breast cancer where all the	7. Left ventricular ejection fraction of 50% or more		(formerly TA371)	15 341 17	27 000 27
		following criteria are met:	8. NOTE: not to be used beyond first disease progression outside the CNS. Do not discontinue if disease progression is within the CNS alone				
			9. No planned treatment breaks of more than 6 weeks beyond the expected cycle length are allowed (to allow any toxicity of current therapy to settle or intercurrent comorbidities to improve).				
			10. will otherwise be used as set out in its Summary of Product Characteristics (SPC).				
			Note: To minimise the risk of errors due to the similarity of the product name Trastuzumab Emtansine (Kadcyla) with that of Trastuzumab the recommendations in the Risk Minimisation Plan educational material from the manufacturer should be followed when prescribing, dispensing and administering the product				
			1. This application is being made by and the first cycle of systemic anti-cancer therapy with trametinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.				
			2. The patient was initially diagnosed with either: - a serous ovarian or peritoneal carcinoma that has recurred with low grade serous histology (invasive micropapillary serous carcinoma or invasive grade 1 serous carcinoma)				
			a secus oranian of peritonian carcinoma trait in a treatment with ow great secus animology (invasive micropapillar) secus occurrence or a started with a serious borderine orania or peritonian carcinoma which has recurred as low-grade serious carcinoma (invasive micropapillar) serious carcinoma or invasive grade 1 carcinoma)				
		F	3. The patient has or had disease which has progressed following at least 1 previous platinum-based chemotherapy regimen.				
		For serous low grade ovarian or peritoneal cancer for disease which has recurred or	4. The patient has not previously received any MEK inhibitors.				
TRAM1	Trametinib		5. Trametinib will be used as monotherapy at a dose of 2 mg daily as part of a 28 day cycle.	No	NHSE Policy:	N/A	08-Nov-23
		based chemotherapy regimen where the	6. The patient has an ECOG performance status of either 0 or 1.	-	URN2253	•	
		following criteria have been met:	6. The patient has an ECO performed valued or included and included an				
			8. A formal medical review as to how tramethin is being tolerated and whether treatment with tramethin is brould continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment.				
			9. Trust policy regarding the use of unlicensed treatments has been followed as this treatment is not licensed in this indication.				
			10. Where a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, the prescribing clinician will complete a treatment break approval form to restart treatment.				
			11. Trametinib is to be otherwise used as set out in its Summary of Product Characteristics.				1

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
TRE1	Treosulfan (Trecondi*) in combination with fludarabine	treatment prior to allogeneic haemopoietic stem cell transplantation for malignant disease in ADUITS for whom a reduced intensity conditioning regimen (such as low dose busulfan with fludarabine) would otherwise be suitable where the following criteria have been	1. This application for treosulfan (as Trecondi*) in combination with fludarabine is being made by and will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy and who has specific expertise in the allogeneic stem cell transplantation of malignant disease.  2. The patient is an adult and the allogeneic stem cell transplantation is for the treatment of malignant disease.  3. This patient is ineligible for high intensity myeloablative therapy and as a consequence a reduced intensity conditioning regimen (such as low dose busulfan plus fludarabine or low dose melphalan plus fludarabine) as treatment prior to allogeneic stem cell transplantation would otherwise be suitable.  4. Treosulfan (as Trecondi*) plus fludarabine will be used as part of the reduced intensity conditioning treatment prior to the allogeneic stem cell transplantation.  Note: Trecondi* is the only licensed formulation of tresosulfan for use in this indication.  5. Treosulfan (as Trecondi*) and fludarabine (including their doses and schedules of administration) will be otherwise used as set out in their respective Summaries of Product Characteristics (SmPCs).	No	TA640	05-Aug-20	03-Nov-20
TRE2	Treosulfan (Trecondi*) in combination with fludarabine	Treosulfan (as Trecondi*) in combination with fludarabine for part of conditioning treatment prior to allogeneic haemopoietic stem cell transplantation for malignant disease in PAEDIATRIC PATENTS OLDER THAN 1 a WONTH AND YOUNGER THAN 18 YEARS for whom a reduced intensity conditioning regimen (such as low dose busulfan with fludarabine) would otherwise be suitable where the following criteria have been met:  There is a separate form TRE1 for treosulfan in combination with fludarabine for part of conditioning treatment prior to allogeneic haemopoietic stem cell transplantation for	1. This application for treosulfan (as Trecondi*) in combination with fludarabine is being made by and will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy and who has specific expertise in the allogeneic stem cell transplantation of malignant disease.  2. The patient is older than 1 month and younger than 18 years patient.  Note: this access to Trecondi* in this indication is a Medicines for Children Policy extension of TA640.  Note: there is a separate application form TR£1 to be used for this indication in adults.  3. Allogeneic stem cell transplantation is for the treatment of malignant disease.  4. This patient is ineligible for high intensity myeloabitive therapy and as a consequence a reduced intensity conditioning regimen (such as low dose busulfan plus fludarabine or low dose melphalan plus fludarabine) as treatment prior to allogeneic stem cell transplantation would otherwise be suitable.  5. Treosulfan (as Trecondi*) plus fludarabine will be used as part of the reduced intensity conditioning treatment prior to the allogeneic stem cell transplantation.  Note: Trecondi* is the only licensed formulation of tresosulfan for use in this indication.  6. The use of treosulfan (as Trecondi*) in combination with fludarabine as a reduced intensity conditioning regimen prior to allogeneic stem cell transplantation has been discussed at a multidisciplinary team (MDT) meeting which must include at least 2 consultants in the subspecialty with active and credible expertise in the relevant field of whom at least one must be a consultant paediatrician. The MDT should include a paediatric pharmacist and other professional groups appropriate to the disease.  7. Treosulfan (as Trecondi*) and fludarabine (including their doses and schedules of administration in this indication) will be otherwise used as set out in their respective Summaries of Product Characteristics (SPCs).	No	TA640	05-Aug-20	09-May-24

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
TRI1_v1.2	Trifluridine plus tipiracil	For patients with either metastatic or locally advanced and inoperable colorectal cancer who have been previously treated with, or are not considered candidates for, available therapies including fluoropyrimidine-based chemotherapy and anti-EGFR-based treatment where the following criteria have been met:	1. This application is both being made by and the first cycle of systemic anti-cancer therapy.  2. The patient has a histologically confirmed diagnosis of adenocarcinoma of the colon or rectum.  3. The patient has either metastatic or locally advanced and inoperable disease.  4. The patient has been previously treated for metastatic disease with, or is not considered a candidate for, fluoropyrimidine-containing chemotherapies which include 5-fluorouracil and/or capecitabine and/or tegafur but not trifluridine (plus tipiracil).  5. The patient has been previously treated with, or is not considered a candidate for, anti-EGFR-containing chemotherapy.  6. The patient has been previously treated with regorafenib or not.  Please tick which option applies to this patient:  - ves, the patient has one previously treated with regorafenib or - on, the patient has not been previously treated with regorafenib or - on, the patient has not been previously treated with regorafenib  7. The patient has not been previously treated with regorafenib  7. The patient has not been previously treated with regorafenib  7. The patient has not been previously treated with regorafenib  7. The patient has not been previously treated with regorafenib  7. The patient has not been previously treated with regorafenib  7. The patient has not been previously treated with regorafenib  7. The patient has not been previously treated with regorafenib  7. The patient has not been previously treated with regorafenib  7. The patient has not been previously treated with regorafenib  7. The patient has not been previously treated with regorafenib  8. The patient has not been previously treated with regorafenib  9. Triffuridine plus tipiracil is to be continued until loss of clinical benefit or unacceptable toxicity or patient choice to stop treatment, whichever is the sooner.  10. A formal medical review as to whether treatment with triffuridine plus tipiracil should continue or not will be scheduled to occur no later than by the end of the 2nd (28-day) c	No	TA405	24-Aug-16	22-Nov-16
TRI2_v1.1	Trifluridine plus tipiracil	For the third or more line of systemic therapy for locally advanced or metastatic adenocarcinoma of the stomach or gastro-oesophageal junction where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-cancer therapy with trifluridine plus tipiracil will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a histologically confirmed diagnosis of adenocarcinoma of the stomach or gastro-oesophageal junction.  3. The patient has been treated with 2 or more systemic therapy regimens for locally advanced or metastatic disease.  4. The patient has an ECOG performance status of 0 or 1.  5. The patient has not been previously treated with trifluridine plus tipiracil.  6. Trifluridine plus tipiracil is not to be used in combination with any other systemic anti-cancer therapy.  7. Trifluridine plus tipiracil is to be continued until loss of clinical benefit or unacceptable toxicity or patient choice to stop treatment, whichever is the sooner.  8. A formal medical review as to whether treatment with trifluridine plus tipiracil should continue or not will be scheduled to occur no later than by the end of the 2nd (28-day) cycle of therapy.  9. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, a treatment break form will be completed to restart treatment.  10. Trifluridine plus tipiracil will be otherwise used as set out in its Summary of Product Characteristics.	No	TA852	14-Dec-22	14-Mar-23

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
TRI3	Trifluridine plus tipiracil in combination with bevacizumab	cancer who have received 2 or more prior anticancer treatment regimens including fluoropyrimidine-, oxaliplatin- and irinotecan-based chemotherapies with without anti-VEGF agents and/or anti-EGFR-based agents where the following criteria have been met:	1. This application is both being made by and the first cycle of systemic anti-cancer therapy with trifluridine plus tipiraciil in combination with bevacizumab will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a histologically confirmed diagnosis of adenocarcinoma of the colon or rectum.  3. The patient has either metastatic disease or locally advanced and inoperable disease.  4. The patient has been previously treated for metastatic or locally advanced and inoperable disease with 2 or more prior anticancer regimens including fluoropyrimidine, oxaliplatin- and innotecan-based chemotherapies. If disease has recurred during or within 6 months after the last administration of neoadjuwant or adjuvant therapy, this can be counted as a prior line of treatment for metastatic or locally advanced and inoperable disease. Note: the regimens of either FOLIRININO or FOLIRININO or FOLIRININO are FOLIRININO are FOLIRININO or FOLIRININO are Folirining Chemotherapy or not.  Please tick which option applies to this patient:  - yes, the patient has been previously treated with an anti-VEGF-containing chemotherapy or not, the patient has not been previously treated with an anti-VEGF-containing chemotherapy or not, the patient has not been previously treated with regorafenib or not, the patient has not been previously treated with regorafenib or not, the patient has not been previously treated with regorafenib or not, the patient has not been previously treated with the folirinine plus tipiracil and at a dose o	No	TA1008	25-Sep-24	24-Dec-24
			14. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, a treatment break approval form will be completed to restart treatment.  15. Both trifluridine plus tipiracil and bevacizumab will be otherwise used as set out in their respective Summaries of Product Characteristics (SPCs).	-			

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
TUC1	<b>Tucatinib</b> in combination with trastuzumab and capecitabine	For treating over-expressed HER2 positive unresectable locally advanced or metastatic breast cancer after 2 or more anti-HER2 treatment regimens where the following criteria have been met:	1. This application for tuactivib in combination with trasturumab and capecitabine for the treatment of unrescrable locally advanced or metastatic breast cancer is being made by and the first cycle of this turativib combination will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has intersectable locally advanced or metastatic breast cancer.  3. The patient has intersectable locally advanced metastatic breast cancer.  4. Confirmation of whether this patient received at REFA targeted enadigivant regimen and if so its nature.  Please tick which option applies to this patient:  1. the patient was treated with a REFA targeted enadigivant regimen which contained trasturumab as the sole HER2-targeted agent.  5. Confirmation of whether the patient received at REFA targeted adjuvant regimen and if so its nature.  Please tick which option applies to this patient:  1. the patient was not treated with a HER2-targeted adjuvant regimen and if so its nature.  Please tick which option applies to this patient:  1. the patient was not treated with a HER2-targeted adjuvant regimen which contained trasturumab and trasturumab and trasturumab and trasturumab and trasturumab and trasturumab and trasturumab and trasturumab.  1. the patient was reteated with a HER2-targeted adjuvant regimen which contained trasturumab as the sole HER2-targeted agent.  1. the patient was reteated with a HER2-targeted adjuvant regimen which contained trasturumab and trasturumab.  2. Confirmation of whether the patient received at HER2-targeted adjuvant regimen which contained trasturumab are manning.  3. Confirmation of whether the patient received at HER2-targeted agine manning the patient received at HER2-targeted agine manning the patient received at HER2-targeted agine manning the patient received at HER2-targeted agine manning the patient received at HER2-targeted regimen for locally advanced/metastatic disease which included both perturumab and trasturumab.	No	TA786	27-Apr-22	26-Јиі-22
			11. The patient has not previously received treatment with tucatinib unless the patient has received tucatinib via a company early access scheme and the patient meets all the other criteria listed here.  12. The patient has not been previously treated with capecitabline in the locally advanced/metastatic disease setting.  13. The status as to the presence of brain metastases/leptomeningeal spread and its symptomatic and treatment status:  • the patient has never had any known brain metastases or leptomeningeal spread and has not received any active treatment for this CNS spread  • the patient has active brain metastases/leptomeningeal spread and has not received any active treatment for this CNS spread  • the patient has been previously treated with CNS radiotherapy/stereotactic radiosurgery/intrathecal chemotherapy and the metastatic CNS disease is stable  • the patient has been previously treated with CNS radiotherapy/stereotactic radiosurgery/intrathecal chemotherapy and the metastatic CNS disease is progressing  14. The patient has an ECOS performance status of 0 or 1.  15. Confirmation of whether the treatment intent for all the treatment period is for this patient to receive trastuzumab via its subcutaneous or intravenous formulations.  1ts strongly received commended by NHS England that the patient is treated with subcutaneous trastuzumab from the start of treatment with tucatinib plus capecitabine. The subcutaneous administration of trastuzumab has obvious benefit for patients and significant service capacity advantages over intravenous administration for providers.  Please mark below whether the treatment intent for all the treatment period with tucatinib in combination with trastuzumab and capecitabine is to use the subcutaneous or the intravenous formulations of trastuzumab:  • subcutaneous trastuzumab is preferred for the entire treatment period  • intravenous trastuzumab is preferred for the entire treatment period  • intravenous trastuzumab is preferred for the entire treatment period  • intraveno				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
VEN1_v1.1	Venetoclax monotherapy	Treatment of chronic lymphatic leukaemii in the ABSENCE of 17p deletion (and absence of TPS3 mutation if tested) wher the following criteria have been met:	7. The patient has never received venetoclax before or has been previously treated with the combination of venetoclax with an anti-CD20 antibody (obinutuzumab or rituximab) or the combination of ibrutinib plus venetoclax in which	No	TA796	15-Jun-22	15-Jul-22

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
VEN2_v1.1	Venetoclax monotherapy	The treatment of previously treated chronic lymphatic leukaemia in the PRESENCE of 17p deletion or TP53 mutation where the following criteria have been met:	1. This application for venetocks plus intusimable is being made by and the first cycle of this systemic anti-cancer therapy.  2. The patient has been diagnosed with chronic lymphatic leukaemia or small lymphocytic lymphoma that requires treatment.  3. The patient has been diagnosed with chronic lymphatic leukaemia or small lymphocytic lymphoma that requires treatment.  4. The prescribing clinician can confirm whether the patient was previously treated with chemoimmunotherapy and if so, then the patient must have had progressive disease.  Please mark below wich applies to this patient:  - the patient has previously been treated with chemoimmunotherapy and had progressive disease on/after such treatment  5. The patient has previously been treated with chemoimmunotherapy and had progressive disease on/after such treatment  5. The patient had progressive disease on or after treatment with a 8 cell receptor pathway inhibitor: a Bruton's tyrosine kinase inhibitor (BTKI e.g., ibrutinib, acalabrutinib) and/or a PI3K inhibitor (PI3KI e.g. idealisib) or has a contraindication to charge on/after a PI3K  - relapse on/after a PI3K  - relapse on/after a PI3K  - relapse on/after a PI3K  - relapse on/after a PI3K  - relapse on/after a PI3K  - relapse on/after a PI3K  - relapse on/after both a BTXI and a PI3KI. Please indicate which:  - a previous lines of treatment  - a previous lines of treatment  - a previous lines of treatment  - a previous lines of treatment  - a previous lines of treatment  - a previous lines of previous treatment  - a relative to the after a previous treatment  - a relative to the after a previous treatment  - a relative to the after a previous treatment  - a relative to the after a previous treatment with venetoclax.  - previous treatment with the combination of venetoclax and riturnish and there was no disease progression whilst on venetoclax  - previous treatment with the combination of venetoclax and dibuntable and there was no disease progression whilst on venetoclax  - previous treatment with the	No	TA796	15-Jun-22	15-Jul-22

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Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
VEN3_v1.7	Venetoclax (in combination with rituximab)	The treatment of previously treated chronic lymphatic leukaemia	1. This agalication for venetociar plus risusmob is being made by and the first cycle of this systemic anti-cancer therapy will be prescribed by a consultant specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has been teasef for 17-95 delition. Please indicate the result of this test below.  3. The patient has been teasef for 17-95 mustion or has not been tested for 17-95 mustion or has not been tested for 17-95 mustion or has not been tested for 17-95 mustion or has not been tested for 17-95 mustion or has not been tested for 17-95 mustion or has not been tested for 17-95 mustion or has not been tested for 17-95 mustion or has not been tested for 17-95 mustion or has not been tested for 17-95 mustion or has not been tested for 17-95 mustion or has not been tested for 17-95 mustion or  No	TA561	27-Feb-19	started	

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
VENS	<b>Venetoclax</b> In combination with obinutuzumab	For the treatment of patients with previously untreated chronic lymphatic leukaemia which has a 17p deletion or TPS3 mutation where the following criteria have been met:	1. This application for venetoclax plus obinutuzumab is being made by and the first cycle of this systemic anti-cancer therapy.  2. The patient has been diagnosed with chronic lymphatic leukaemia (CLL) or small lymphocytic lymphoma (SLL).  3. The patient has been tested for 17p deletion and for TP53 mutation and the results are positive for 17p deletion or TP53 mutation or both.  Please indicate the result of these tests below:  - Positive for 17p deletion and negative for TP53 mutation or - Positive for 17p deletion and positive for TP53 mutation or - Positive for 15p deletion and positive for TP53 mutation or - Positive for both 17p deletion and TP53 mutation.  4. The patient has symptomatic disease which requires systemic therapy.  5. The patient has not received any previous systemic therapy for CLL/SLL.  6. The patient has a performance status of 0 or 1 or 2.  7. Venetoclax will be given in combination with obinutuzumab and that the venetoclax dose titration schedule is planned to commence on cycle 1 day 22 and be completed on cycle 2 day 28.  8. All of the following for the prevention and treatment of tumour lysis syndrome: - that the patient has been prospectively assessed for the risk of the development of tumour lysis syndrome (TLS) with venetoclax - that the praintent TLS risk militagion strategies have been put in place as outlined in the updated venetoclax Summary of Product Characteristics. See https://www.medicines.psu.psu.psu.psu.psu.psu.psu.psu.psu.ps	-	TA663	09-Dec-20	09-Mar-21
			9. The patient has been assessed specifically for potential drug interactions with venetoclax.  10. The maximum treatment duration of venetoclax in this indication is until day 28 of the 12th cycle of treatment i.e. the maximum duration of venetoclax treatment is for 45 weeks, consisting of 1 week from cycle 1 day 22 followed by 11 cycles of 4-weekly cycles of venetoclax in cycles 2-12.  11. The treatment duration of obinutuzumab is for a maximum of 6 cycles of obinutuzumab.  12. Venetoclax is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment or for the maximum treatment duration of 12 cycles (as measured above), whichever of these events is the sooner.  13. A formal medical review as to whether treatment with venetoclax in combination with obinutuzumab should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment.  14. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment, including as appropriate if the patient had an extended break on account of Covid-19.  15. Venetoclax and obinutuzumab will be otherwise used as set out in their respective Summary of Product Characteristics (SPCs).				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
VENG	<b>Venetoclax</b> in combination with obinutuzumab	For the treatment of patients with previously untreated chronic lymphatic leukaemia in whom chemotherapy with the combinations of either FCR or BR. would otherwise have been UNSUITABLE where the following criteria have been met:	1. This application for venetoclax plus obinutuzumab is being made by and the first cycle of this systemic anti-cancer therapy.  2. The patient has been diagnosed with chronic lymphatic leukaemia (CLL) or small lymphocytic lymphoma (SLL).  3. The patient has been tested for 17p deletion and the result is negative.  4. The patient has been tested for 17p3 mutation and the result is negative.  5. The patient has symptomatic disease which requires systemic therapy.  6. The patient has not received any previous systemic therapy for CLL/SLL.  7. The patient has not received any previous systemic therapy for CLL/SLL.  7. The patient has not received any previous systemic therapy for CLL/SLL.  7. The patient has a performance status of 0 or 1 or 2.  8. In the absence of this venetoclax plus obinutuzumab treatment option, the patient would otherwise have been considered to have been UNSUITABLE for treatment with the combination of fludarabine, cyclophosphamide and rituximab (FCR) or the combination of bendamustine and rituximab (BR).  9. Venetoclax will be given in combination with obinutuzumab and that the venetoclax dose titration schedule is planned to commence on cycle 1 day 22 and be completed on cycle 2 day 28.  10. All of the following for the prevention and treatment of tumour lysis syndrome:  1. That paper has been prospectively assessed for the risk of the development of tumour lysis syndrome (TLS) with venetoclax  1. That appropriate TLS risk mitigation strategies have been put in place as outlined in the updated venetoclax Summary of Product Characteristics  1. That there is a robust system in place for measuring appropriate blood chemistries both at the specified timings of blood chemistries according to TLS risk status and at the venetoclax dose levels described in Section 4.2 Table 3 of the Summary of Product Characteristics. See https://www.medicines.org.uk/em/cmedicine/32650 or https://products.mhra.gov.uk/substance/substance-VENETOCLAX  1. That there is a robust system in place for measuring appropria		TA663	09-Dec-21	09-Mar-21
			11. The patient has been assessed specifically for potential drug interactions with venetoclax.  12. The maximum treatment duration of venetoclax in this indication is until day 28 of the 12th cycle of treatment i.e. the maximum duration of venetoclax treatment is for 45 weeks, consisting of 1 week from cycle 1 day 22 followed by 11 cycles of 4-weekly cycles of venetoclax in cycles 2-12.  13. The treatment duration of obinutuzumab is for a maximum of 6 cycles of obinutuzumab.  14. Venetoclax is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment or for the maximum treatment duration of 12 cycles (as measured above), whichever of these events is the sooner.  15. A formal medical review as to whether treatment with venetoclax in combination with obinutuzumab should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment.  16. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment, including as appropriate if the patient had an extended break on account of Covid-19.  17. Venetoclax and obinutuzumab will be otherwise used as set out in their respective Summary of Product Characteristics (SPCs).	-			

26-Nov-2025

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
VEN8	Venetoclax in combination with azacitidine	For untreated adult acute myeloid leukaemia in patients unsuitable for intensive chemotherapy where the following criteria have been met:	1. This application is bring made by and the first cycle of sptemic anti-cancer therapy.  2. The patient has newly diagnosed acute myeloid leukaemia (AML).  3. The patient has newly diagnosed acute myeloid leukaemia (AML).  3. The patient has newly diagnosed acute myeloid leukaemia (AML).  3. The patient has newly diagnosed acute myeloid leukaemia (AML).  3. The patient has newly diagnosed acute myeloid leukaemia (AML).  3. The patient has newly diagnosed acute myeloid leukaemia (AML).  4. The patient has performed.  4. The patient has performed.  4. The patient has previously untreated de novo AML or previously untreated secondary AML.  4. The patient has previously untreated de novo AML or previously untreated secondary AML.  4. The patient has previously untreated de novo AML or previously untreated secondary AML.  5. The most remet home marrow blast count is:  4. The patient has previously untreated de novo AML or previously untreated secondary AML.  5. The most remet home marrow blast count is:  4. The patient has previously untreated de novo AML or previously untreated secondary AML.  5. The most remet home marrow blast count is:  4. The patient has previously untreated de novo AML or previously untreated secondary AML.  5. The most remet home marrow blast count is:  4. The patient has previously untreated de novo AML or previously untreated secondary AML.  5. The most remet home marrow blast count is:  4. The patient has been prospectively assessed for the fixed of the development of the more previously untreated secondary is unsuitable for intensive chemotherapy:  4. The patient has been prospectively assessed for the fix of the development of fixed previously untreated pre	No	TA765	02-feb-22	03-May-22

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
VEN9	Venetoclax in combination with low dose cytarabine	For previously untreated adult acute myeloid leukaemia in patients unsuitable for intensive chemotherapy and who have a bone marrow blast count -350% where the following criteria have been met:	1. This application is being made by and the first cycle of systemic anti-carect therapy.  2. The patient has newly diagnosed acute myeloid leukaemia (ANL).  3. The patient has newly diagnosed acute myeloid leukaemia (ANL).  3. The patient has newly diagnosed acute myeloid leukaemia (ANL).  3. The patient has newly diagnosed acute myeloid leukaemia (ANL).  3. The patient has had; having molecular analysis performed.  4. Plasse mank below the somatic mutation found:  - not system with below the somatic mutation found:  - not system with a system of the patient of the pat	No	TA787	27-Apr-22	26-Jul-22

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
VI52	Vismodegib	For patients with multiple basal cell carcinomas (BCC) in adults where the	1. This application is being made by and the first cycle of systemic anti-cancer therapy with vismodegib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy  2. The patient has either (tick as appropriate):  - Gorlin syndrome with non-locally advanced, non-metastatic multiple basal cell carcinomas (BCC) (2-6) clinically evident at the point of decision to treat BCCs of which 3 are at least 5mm or  - Non-locally advanced, non-metastatic multiple BCC (2-6) clinically evident at the point of decision to treat BCCs of which 3 are at least 5mm AND are appropriate for surgery i.e. surgically eligible tumours.  3. The patient has at least 6 operable clinically evident non-locally advanced, non-metastatic BCC with surgically eligible tumours of 3 lesions of at least 5mm diameter, of which at least 1 is histopathologically confirmed.  4. The patient has been assessed and vismodegib recommended by a specialised skin cancer or head and neck multidisciplinary team.  5. The patient has been assessed and vismodegib recommended by a specialised skin cancer or head and neck multidisciplinary team.  6. The patient has been explained and agreed with the patient before the treatment is started.  8. Vismodegib will be prescribed at a dose of 150mg daily taken once daily OR on an intermittent schedule, until disease progression or adverse effects which necessitate stopping.  Please note which treatment schedule will be used (tick box):  - Continuous therapy or  - A 72 week period of: vismodegib 12 weeks; off treatment 8 weeks; vismodegib 12 weeks; off treatment 8 weeks; vismodegib 8 weeks:  - A 72 week period of: vismodegib 24 weeks; off treatment 8 weeks; vismodegib 8 weeks:	No.	NHSE Policy:	n/a	started
V132	Tismodegis	following criteria have been met:	*Reference: Dreno, B., Kunstfeld, R., Hauschild, A., Fosko, S., Zloty, D., Labeille, B., Grob, J-J. et al. (2017) Two intermittent vismodegib dosing regimens in patients with multiple basal-cell carcinomas (MIKIE): a randomised regimen-controlled, double-blind, phase 2 trial. The Lancet Oncology 18:404-12.		210504P	11/4	14-301-21
			9. The patient is either male or female  10. The prescibing clinician understands that vismodegib must not be used during pregnancy and female and male patients will be counselled as describe below.  10. The prescibing clinician understands that vismodegib must not be used during pregnancy and female and male patients will be counselled as describe below.  10. The patient has been counselled about the adverse use of vismodegib in pregnancy AND, lif a woman of child-bearing potential, has been advised that she should use two forms of contraception (including one highly effective method and one barrier) during vismodegib therapy and for 24 months after the final dose, AND has had a negative medically supervised pregnancy test within the past seven days.  10. Counselling for male patients:  11. The patient has been counselled about the adverse use of vismodegib in relation to pregnancy and has been advised that he should always use a condom (with spermicide if available), during vismodegib therapy and for 2 months after the final dose.				
			11. This application is for an adult patients and vismodegib will not be used in children and adolescents aged below 18 years.  12. Trust policy regarding the use of unlicensed treatments has been followed as vismodegib and the recommended intermittent schedules are not licensed in this indication.				
			12. Tota pointy regarding the set of unitensed treatments has been followed as visinousgo and the recommended intermediate intermediate intermediate intermediate intermediate intermediate intermediate intermediate intermediate.  13. Where a treatment break of more than 6 weeks beyond the expected 4-week (yel length is needed, I will complete a treatment break approval form to restart treatment, including indicating as appropriate if the patient had an extended break because of COVID 19.  14. Visinodesjib will oftherwise be used as set out its Summary of Product Characteristics				

26-Nov-2025

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ZAN1	Zanubrutinib	Zanubrutinib monotherapy for the treatment of patients with previously treated Waldenstrom's macroglobulinaemia and who would otherwise be next treated with bendamustine plus rituximab where the following criteria have been met:	1. This application is being made by and the first cycle of this systemic anti-cancer therapy with zanubrutinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has been previously diagnosed with Waldenstrom's macroglobulinaemia.  3. The patient has symptomatic disease which requires systemic therapy.  4. The patient has been previously treated with at least 1 prior systemic therapy for Waldenstrom's macroglobulinaemia.  Note: NICE could not recommend the use of zanubrutinib in treatment-naïve patients in whom chemo-immunotherapy is unsuitable as the company did not submit evidence for the clinical and cost effectiveness of zanubrutinib in this patient group.  5. In the absence of this access to zanubrutinib, the patient would otherwise be next treated with the combination of bendamustine and rituximab.  Note: the only previously treated patient group for which NICE concluded that zanubrutinib was clinically and cost effective was in those patients who would otherwise be next treated with the combination of dexamethasone, rituaimab and cyclophosphamide or any other therapies.  6. The patient is treatment naïve to a Bruton's kinase inhibitor or the patient has been previously commenced on ibrutinib for previously treated Waldenstrom's macroglobulinaemia and all other treatment criteria on this from are fulfilled or the patient has been previously commenced on ibrutinib for previously treated Waldenstrom's macroglobulinaemia and the ibrutinib has had to be discontinued solely as a consequence of observable part previously commenced zanubrutinib in a the manufacturer's (BeiGene) early access scheme for previously treated Waldenstrom's macroglobulinaemia and the ibrutinib has had to be discontinued solely as a consequence of dose-limiting toxicity and in the clear absence of disease progression  1. The patient previously commenced zanubrutinib via the manufacturer's (BeiGene) early access scheme for previously treated Walde		TA833	19-Oct-22	17-Jan-23
			7. The patient has an ECOG performance status of 0 or 1 or 2.  8. The use of zanubrutinib in this indication will be as monotherapy.  9. The prescribing clinician is aware that zanubrutinib has clinically significant drug interactions with CYP3A inhibitors and inducers as described in zanubrutinib's Summary of Product Characteristics.  10. Zanubrutinib is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment, whichever is the sooner.  11. A formal medical review as to whether treatment with zanubrutinib should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment.  12. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, a treatment break approval form will be completed to restart treatment, including as appropriate if the patient had an extended break on account of Covid-19.  13. Zanubrutinib will be otherwise used as set out in its Summary of Product Characteristics (SPC).				
ZAN2_v1.0	<b>Zanubrutinib</b> monotherapy	For the treatment of patients with previously untreated chronic lymphatic leukaemia which has a 17p deletion or TPS3 mutation where the following criteria have been met:	1. This application for zanubrutninb is being made by and the first cycle of this systemic anti-cancer therapy.  2. The patient has been diagnosed with chronic lymphatic leukaemia (CLL) or small lymphocytic lymphoma (SLL).  3. The patient has been diagnosed with chronic lymphatic leukaemia (CLL) or small lymphocytic lymphoma (SLL).  3. The patient has been tested for 17p deletion and for TP53 mutation and the results are positive for 17p deletion or or TP53 mutation or both.  Please indicate the result of these tests below:  - negative for 17p deletion and positive for TP53 mutation or - negative for 17p deletion and positive for TP53 mutation or - positive for 10p deletion and positive for TP53 mutation or - positive for both 17p deletion and TP53 mutation.  4. The patient has symptomatic disease which requires systemic therapy.  5. The patient has not received any previous systemic therapy for CLL/SLL unless 1st line zanubrutinib was previously commenced via a BeiGene early access scheme or 1st line acalabrutinib or 1st line ibrutinib has had to be stopped due to dose-limiting toxicity and in the clear absence of disease progression.  Please mark which of the 4 scenarios below applies to this patient: - the patient previously commenced 1st line acalabrutinib as a BeiGene early access scheme and all other treatment criteria on this form are fulfilled or - the patient previously commenced 1st line acalabrutinib and the acalabrutinib has had to be stopped solely due to dose-limiting toxicity and in the clear absence of disease progression - the patient previously commenced 1st line acalabrutinib and the ibrutinib has had to be stopped solely due to dose-limiting toxicity and in the clear absence of disease progression - the patient previously commenced 1st line acalabrutinib and the ibrutinib has had to be stopped solely due to dose-limiting toxicity and in the clear absence of disease progression - the patient previously commenced 1st line acalabrutinib has bad to be stopped solely due to dose-limiting toxicity	No	TA931	22-Nov-23	20-Feb-24

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ZAN3_v1.0	<b>Zanubrutinib</b> monotherapy	For the treatment of patients with previously untreated chronic lymphatic leukaemia which does not have a 170 deletion or a 1753 mutation and in whom chemotherapy with FCR or BR is unsuitable where the following criteria have been met:	1. This application for zanubrutinib is being made by and the first cycle of this systemic anti-cancer therapy with zanubrutinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has been diagnosed with chronic lymphatic leukaemia (CLL) or small lymphocytic lymphoma (SLL).  3. The patient has been tested for 17p deletion and the result is negative.  4. The patient has been tested for 17p deletion and the result is negative.  5. The patient has symptomatic disease which requires systemic therapy.  6. In the absence of this zanubrutinib treatment option, the patient would otherwise have been considered as UNSUITABLE for treatment with the combination of fludarabine, cyclophosphamide and rituximab (FCR) or the combination of bendamustine and rituximab (BR).  Note: NICE's assessment of the clinical and cost effectiveness of 1st line zanubrutinib resulted in a positive recommendation for zanubrutinib to be an option in those places in the treatment pathway which have current recommendations for use of a BTK inhibitor as monotherapy.  7. The patient has not received any previous systemic therapy for CLL/SLL unless 1st line zanubrutinib was previously commenced via a BeiGene early access scheme or 1st line acalabrutinib has had to be stopped solely due to dose-limiting toxicity and in the clear absence of disease progression.  Please mark which of the 3 scenarios below applies to this patient:  - the patient has not received any systemic therapy for CLL/SLL i.e. is completely treatment-naive or  - the patient previously commenced 1st line acalabrutinib was beingen early access scheme and all other treatment criteria on this form are fulfilled.  - the patient previously commenced 1st line acalabrutinib and the acalabrutinib has had to be stopped solely due to dose-limiting toxicity and in the clear absence of disease progression  8. The patient has an ECOS performance status of 0 or 1 or 2.  9. Use of zanubrutinib in this indica	indication  No		22-Nov-23	started 20-Feb-24
			10. The prescribing clinician is aware that zanubrutinib has clinically significant interactions with cytochrome P450 enzyme 3A (CYP3A) inhibitors and inducers as described in zanubrutinib's Summary of Product Characteristics.  11. Zanubrutinib is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment, whichever is the sooner.  12. A formal medical review as to whether treatment with zanubrutinib should continue or not will be scheduled to occur at least by the end of the first 8 weeks of treatment.  13. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, a treatment break approval form will be completed to restart treatment.  14. Zanubrutinib will be otherwise used as set out in its Summary of Product Characteristics (SPC).				
ZAN4_v1.0	<b>Zanubrutinib</b> monotherapy	For the treatment of patients with previously treated chronic lymphatic leukaemia where the following criteria have been met:	1. This application for zanubrutnib is being made by and the first cycle of this systemic anti-cancer therapy.  2. The patient has been previously diagnosed with chronic lymphatic leukaemia (CLL) or small lymphocytic lymphoma (SLL).  3. The patient has been tested for 17p deletion and for TP53 mutation and the results are as shown below:  - negative for both 17p deletion and negative for TP53 mutation or - negative for 17p deletion and positive for 17p3 mutation or - negative for 17p deletion and positive for TP53 mutation or - negative for 17p deletion and positive for 17p3 mutation or - positive for 17p deletion and positive for TP53 mutation or - positive for 17p deletion and positive for TP53 mutation or - positive for 10p4 17p deletion and positive for TP53 mutation or - positive for both 17p deletion and positive for TP53 mutation or - positive for both 17p deletion and positive for TP53 mutation or - positive for both 17p deletion and positive for TP53 mutation - positive for 10p4 positive for both 17p deletion and positive for TP53 mutation or - positive for 10p4 positive for both 17p deletion and positive for TP53 mutation or - positive for 10p4 positive for both 17p deletion and positive for TP53 mutation or - positive for 10p4 positive for 1	No	TA931	22-Nov-23	20-Feb-24
			7. The patient has an ECOG performance status of 0 or 1 or 2.  8. Use of zanubrutinib in this indication will be as monotherapy.  Note: zanubrutinib is not licensed in CLL to be used in combination with any other agent.  9. The prescribing clinician is aware that zanubrutinib has clinically significant interactions with cytochrome P450 enzyme 3A (CYP3A) inhibitors and inducers as described in zanubrutinib's Summary of Product Characteristics.  10. Zanubrutinib is to be continued until disease progression or unacceptable toxicity or patient choice to stop treatment, whichever is the sooner.  11. A formal medical review as to whether treatment with zanubrutinib should continue or not will be scheduled to occur at least by the end of the first 12 weeks of treatment.  12. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, a treatment break approval form will be completed to restart treatment.  13. Zanubrutinib will be otherwise used as set out in its Summary of Product Characteristics (SPC).				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ZAN5	Zanubrutinib	Zanubrutinib monotherapy for the treatment of patients with marginal zone lymphoma treated with at least 1 prior anti-CD20-based therapy where the following criteria have been met:	1. This application for zanubrutinib is being made by and the first cycle of this systemic anti-cancer therapy with zanubrutinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a confirmed histological diagnosis of marginal zone lymphoma (MZL).  3. The patient has been previously treated with at least 1 prior anti-CD20- based regimen for MZL.  Please mark below how many lines of systemic therapy the patient has received:  - the patient has had 2 prior lines of systemic therapy of which at least one line of treatment contained an anti-CD20 agent or  - the patient has had 2 prior lines of systemic therapy of which at least one line of treatment contained an anti-CD20 agent or  - the patient has had 3 prior lines of systemic therapy of which at least one line of treatment contained an anti-CD20 agent or  - the patient has had 3 prior lines of systemic therapy of which at least one line of treatment contained an anti-CD20 agent or  - the patient has had 3 prior lines of systemic therapy of which at least one line of treatment contained an anti-CD20 agent or  - the patient has had 3 prior lines of systemic therapy of which at least one line of treatment contained an anti-CD20 agent or  - the patient has had 3 prior lines of systemic therapy of which at least one line of treatment contained an anti-CD20 agent or  - the patient has had 3 prior lines of systemic therapy of which at least one line of treatment contained an anti-CD20 agent or  - the patient has had 4 or more prior lines of systemic therapy of which at least one line of treatment contained an anti-CD20 agent or  - the patient has had 2 prior lines of systemic therapy of which at least one line of treatment contained an anti-CD20 agent or  - the patient has had 2 prior lines of systemic therapy of which at least one line of treatment contained an anti-CD20 agent or  - the patient has had 2 prior lines of systemic therapy of which at least one line of treatment	No	TA1001	Guidance  04-Sep-24	_
			11. When a treatment break of more than 6 weeks beyond the expected 4-weekly cycle length is needed, I will complete a treatment break approval form to restart treatment.  12. Zanubrutinib will be otherwise used as set out in its Summary of Product Characteristics (SPC).				

Blueteq Form ref:	Drug	NICE Approved Indication	Blueteq Approval Criteria	Previous CDF drug/ indication	TA	Date of Final NICE Guidance	Date baseline funding started
ZANG	Zanubrutinib		1. This application is being made by and the first cycle of systemic anti-cancer therapy with zanubrutinib will be prescribed by a consultant specialist specifically trained and accredited in the use of systemic anti-cancer therapy.  2. The patient has a confirmed histopathological diagnosis of mantle cell lymphoma.  3. The patient has previously been treated with one and only one prior line of rituximab-containing chemotherapy.  Note: Patients treated with more than 1 line of prior therapy are not eligible for treatment with zanubrutinib.  4. The presence of relapsed/refractory mantle cell lymphoma with documented progression of disease during or following rituximab-containing 1st line systemic therapy.  5. The patient has never received any prior therapy with a BTK inhibitor (ibrutinib or another BTK inhibitor) unless the patient has either received zanubrutinib via a company early access scheme and all other treatment criteria on this form apply or the patient has suffered unacceptable toxicity on therapy with ibrutinib without any evidence of disease progression and is transferring to treatment with zanubrutinib.  Please enter below which of these scenarios applies to this patient:  - the patient is treatment-naive to a BTK inhibitor or  - the patient has received zanubrutinib via a company early access scheme and all other treatment criteria on this form apply or  - the patient has been receiving line therapy with ibrutinib but has suffered unacceptable toxicity without any evidence of disease progression and is transferring to treatment with zanubrutinib.	No	TA1081	10-Jul-25	09-Aug-25
			6. Zanubrutinib is to be used as a single agent. 7. Zanubrutinib is to be used as a single agent. 8. The patient's ECOG performance status is 0 or 1 or 2. 9. The patient is not on concurrent therapy with warfain. 10. The prescribing clinician I am aware that zanubrutinib has clinically significant interactions with cytochrome P450 enzyme 3A (CYP3A) inhibitors and inducers as described in zanubrutinib's Summary of Product Characteristics. 11. When a treatment break of more than 6 weeks beyond the expected cycle length occurs, the prescribing clinician will complete a treatment break approval form to restart treatment. 12. Zanubrutinib will be otherwise used as set out in its Summary of Product Characteristics (SPC).				

#### Section C. Interim Systemic Anti-Cancer Therapy (SACT) treatment change options introduced during the COVID-19 pandemic.

To support the response to the COVID pandemic, NHS England and NICE published a guideline on the delivery of SACT (NICE NG161) and commissioned a list of 'COVID-friendly' interim cancer treatment options. These allowed clinicians to treat patients with less toxic therapies compared to standard treatment and could be given at home.

These arrangements maximised the safety of cancer patients due to start or on chemotherapy during the pandemic response, whilst also preserving efficacy, as well as making the best use of NHS resources (service capacity) and protecting staff from infection and lightening the burden on hospitals, critical during the pandemic response.

Funding for the Interim COVID treatments was provided from the start of the pandemic until the end of 2022/23. The number of Interim options available has decreased over time as indications were removed either because they had been superseded by NICE guidance or the need for the flexibility, they provided during the pandemic has reduced and clinicians have reverted to standard commissioned treatment options.

From 1st April 2023 four options have been retained until the agreed exit strategy for those indications is complete i.e., a decision from NICE which supersedes the COVID-friendly interim option or completion of assessment of a Clinical Policy application by the NHS England Specialised Services Clinical Panel. The options will be removed from this list when the final commissioning position is known or sooner if there is no longer a clinical need to retain these options.

Blueteq Form ref:	Drug Indication	Criteria for use	Date form made available	NICE Guideline	Comment
NIV13CV_v1.1	As 2nd line or subsequent line for malignant pleural and pe mesothelioma which has pro during/after 1st line chemoth pemetrexed- and platinum chemotherapy where the for criteria have been me	7. In terms of previous systemic therapy the patient has only been treated with cytotoxic chemotherapy (which has included first-line pemetrexed and platinum-based combination chemotherapy) and thus this application for nivolumab monotherapy is for second or a subsequently line of systemic treatment.  8. The patient started 1st line chemotherapy on or before 14th July 2022, i.e. the date until which the only first line option available was chemotherapy.	03-Aug-20	NG161	NICE approved nivolumab plus ipilimumab as a first line immunotherapy option in mesothelioma on 14 July 2022 (see NICE 101609). Therefore, the option to give nivolumab monotherapy instead of second-line chemotherapy to reduce risk of immunosuppression only remains in place for patients who started first-line chemotherapy on or before 14 July 2022, when the only first-line option available was chemotherapy.

#### **Version Control**

Version No.	Date published	Author(s)	Revision summary
version No.	Date published	Addibit(s)	nevision summary
0.1	n/a	D Thomson; P Clark	Initial draft of new CDF list, based on pre-existing national CDF list but updated for changes to the CDF, for review.
1.0	29-Jul-16	D Thomson: P Clark	Final version of new CDF list
1.1	09-Aug-16	P Clark	New addition to CDF list
1.2	18-Aug-16	D Thomson; P Clark	New addition to CDF list and revision of criteria for a number of existing drugs
1.3	24-Aug-16	D Thomson; P Clark	Removal of one drug/indication for baseline funding and date for baseline funding added for existing drugs.
1.4	02-Sep-16	D Thomson; P Clark	Update to Radium criteria and timeline following publication of NICE FAD
1.5	20-Sep-16	D Thomson: P Clark	Removal of two drugs/indications for baseline funding
1.6	27-Sep-16	D Thomson: P Clark	Removal of two drug indications
1.7	04-Oct-16	D Thomson; P Clark	Addition of new CDF drug and date for baseline funding added for existing drugs
1.8	21-Oct-16	D Thomson; P Clark	New addition to CDF list
1.9	25-Oct-16	D Thomson: P Clark	Removal of one drug/indication for baseline funding.
1.10	03-Nov-16	D Thomson: P Clark	Update to eribulin following publication of NICE FAD
1.11	10-Nov-16	D Thomson; P Clark	Update to everolimus following publication of NICE FAD; update to section B - "NICE approved and baseline funded drugs/indications from 1st April 2016"
1.12	17-Nov-16	D Thomson; P Clark	Two new addition to CDF list and update to dasatinib criteria following publication of NICE FAD
1.13	23-Nov-16	D Thomson; P Clark	New addition to CDF list, removal of two drugs/indications for baseline funding and update to Nivolumab timeline following publication of final guidance
1.14	02-Dec-16	D Thomson: P Clark	New addition to CDF list (PEMB1 v1.0); update to neoadjuvant pertuzumab (PER2) criteria.
1.15	12-Dec-16	D Thomson: P Clark	New addition to CDF list (IBR3 v1.0); update to ibrutinib in pretreated CLL (IBR1) criteria.
1.16	21-Dec-16	D Thomson: P Clark	Removal of two drugs/indications for baseline funding; update of five timelines following publication of final NICE guidance; update to pembrolizumab criteria.
1.17	23-Dec-16	D Thomson: P Clark	Removal of one drug/indication for baseline funding; update to pertuzumab criteria
1.18	28-Dec-16	D Thomson: P Clark	Removal of three drugs and indications for baseline funding; removal of pegaspargase.
1.19	12-Jan-17	D Thomson; P Clark	Update to everolimus (RCC) following publication of NICE FAD; update to two timelines following publication of final NICE guidance; update to radium 223 criteria in section B
1,20	10-Feb-17	D Thomson: P Clark	Update to section B - "NICE approved and baseline funded drugs/indications from 1st April 2016"; update of 2 timelines following publication of final NICE guidance; update to ponatinib following ACD
1.21	02-Mar-17	D Thomson: P Clark	Updates to section A - CET1, CET4, PAN3, PAN1. Updates to section B - Ipillinumab + Nivolumab, Dabrafenib + Trametinib
1,22	21-Mar-17	D Thomson: P Clark	Removal of 5 drugs/indications for routine funding and addition to section B. Update to Ipilimumab + Nivolumab criteria.
1.23	11-Apr-17	D Thomson: P Clark	Removal of 1 drugs/indications for routine funding.
1,24	27-Apr-17	D Thomson: P Clark	Removal of 2 drug/indications for routine funding and update to section B. Addition of two drug/indications following publication
1,25	28-Apr-17	D Thomson: P Clark	Following publication of ponatinity in CML FAD - incorporation of 2 previous separate sets of criteria into a single set
1.26	02-May-17	D Thomson: P Clark	Replacement of current criteria for brentuximab in HD with new criteria following publication of NICE FAD and update to blimautmomab in children criteria
1.27	12-May-17	D Thomson: P Clark	Addition of 2 CDF drue/indications and updated of 1 CDF drue/indication following publication of FAD
1.28	31-May-17	D Thomson; P Clark	Removal of 1 drug/indication for routine funding and 1 new drug/indication addition following publication of the FAD
1,29	02-Jun-17	D Thomson: P Clark	2 new drug/indications following publication of FAD
1.30	09-Jun-17	D Thomson; P Clark	3 new drug/indications following publication of 2 FADs; update to existing criteria
1.31	15-Jun-17	B Groves: P Clark	Revision to I drug/indication following publication of FAD
1.32	30-Jun-17	D Thomson: B Groves	Revision to 1 drug/indication in CDF / two drugs in 4 indications moved from CDF to routine commissioning
1.33	10-Jul-17	P Clark: B Groves	nersion of a rung minimateur more rywording in minimateurs more rouncer for outline commissioning.  I new drug/indication following publication of FAD.
1.34	24-Jul-17	P Clark: D Thomson: B Groves	The wing/indication; two drugs entering baseline commissioning, update to OLA2 v1.1 interim funding status
1.35	04-Aug-17	P Clark; D Thomson; B Groves	1 new drug/indication for interim funding before moving into routine commissioning
1.36	08-Aug-17	P Clark; D Thomson; B Groves	1 drug/indication revised and 1 new drug indication added
1.37	10-Aug-17	P Clark; D Thomson; B Groves	and the state of t
1.38	24-Aug-17	P Clark; D Thomson; B Groves	I indication deleted and relaised with updated and separate child and adult treatment criteria. Removal of 1 drug-findication for routine fundine and update to section 8: 2 drues 'available to new patients' status updated
			I minication deleted and replaced with updated and separate round and adout retarment increase, kentowal or 1 original cation for fourther formation and updated on separate round and adout retarment increase, kentowal or 1 original cation would not originate to new patients status updated and separate round and adout retarment increase.  I indication moved into routine commissioning, 1 indication updated to reflect notice period for registering new patients.
1.39	31-Aug-17	D Thomson; B Groves	a micration move micro occurrence of the control of the control occurrence oc
1.40	06-Sep-17	D Thomson; B Groves	z micrations updated to therefore the control to th
1.41	08-Sep-17	P Clark; D Thomson; B Groves	I new dug in 2 indications adver, a sexing indication updated to renect expected entry into rounine commissioning  11 indications moved from CDF to routine commissioning
1.42	26-Sep-17	P Clark; D Thomson; B Groves	11 indications indiced from CDF to footine Commissioning 1 drug/indication added
1.43	28-Sep-17	P Clark; D Thomson; B Groves	1 trag/motation accord 1 drug/motation removed; 2 new CDF indications added
1.44	05-Oct-17	P Clark; D Thomson; B Groves	
1.45	12-Oct-17	P Clark; D Thomson	1 drug/indication revised following interim funding
1.46	13-Oct-17	P Clark; D Thomson	1 new drug/indication entering CDF
1.47	17-Oct-17	P Clark; D Thomson; B Groves	2 drugs/indications moving from CDF to routine commissioning
1.48	01-Nov-17	P Clark; D Thomson; B Groves	1 drug/indication criteria updated
1.49	05-Nov-17	P Clark; D Thomson; B Groves	1 drug/indication criteria removed
1.50	08-Nov-17	P Clark; D Thomson; B Groves	1 drug/indication moved from CDF into routine commissioning

#### Version Control(Cont)

Version No.	Date published	Author(s)	Revision summary
1.51	16-Nov-17	P Clark; D Thomson; B Groves	2 new drug/indications added following publication of FAD
1.52	22-Nov-17	P Clark: D Thomson: B Groves	Notice of removal for 1 drug/indication; treatment criteria clarified for 1 drug/indication; 2 drug/indication titles amended
1.53	05-Dec-17	P Clark; D Thomson; B Groves	2 drugs/indications moved into routine commissioning;
1.54	07-Dec-17	P Clark; D Thomson; B Groves	1 drug/indication with interim funding
1.55	08-Dec-17	P Clark; D Thomson; B Groves	1 drug/indication with interim funding
1.56	14-Dec-17	P Clark; D Thomson; B Groves	1 drug/indication split into two indications; 2 drugs/indication updated with dates for expected entry into routine commissioning
1.57	19-Dec-17	P Clark; D Thomson; B Groves	1 new CDF drug/indication; notice given for 2 drugs/indications attracting interim funding which will move into rountine commissioning in 90-days; 4 updates to criteria (1 CDF, 3 routine)
1.58	02-Jan-18	P Clark; D Thomson	2 drug/indications moving from CDF to routine commissioning; 4 updates to criteria (1CDF, 3 routine); 1 update to IFA section
1.59	17-Jan-18	P Clark: B Groves	1 drug/indication added to the CDF; 1 drug/indication updated
1.60	18-Jan-18	P Clark; D Thomson; B Groves	1 drug/indication updated
1.61	22-Jan-18	B Groves	1 drug/indication delisted
1.62	01-Feb-18	B Groves	3 drugs for 4 indications upated following NICE final guidance
1.63	09-Feb-18	P Clark; D Thomson; B Groves	1 drug/indication for routine commissioning
1.64	12-Feb-18	P Clark; D Thomson; B Groves	1 drug/indication for routine commissioning
1.65	15-Feb-18	P Clark; D Thomson; B Groves	3 drug/indication for routine commissioning which will receive interim CDF funding; 1 drug/indication moved into routine commissioning; 1 drug/indication with updated treatment criteria
1.66	21-Feb-18	B Groves	2 drug/indications updated
1.67	01-Mar-18	P Clark; D Thomson; D Dwyer	1 drug/indication added to the CDF; 1 drug/ indication for routine commissioning which will receive interim CDF funding; 3 drug/indications with updated treatment criteria
1.68	07-Mar-18	D Thomson; D Dwyer	1 indication moved into routine commissioning
1.69	16-Mar-18	P Clark; D Thomson; D Dwyer	1 drug/indication added to the CDF
1.70	20-Mar-18	D Thomson; D Dwyer	2 drugs/indications moved into routine commissioning
1.71	21-Mar-18	D Thomson; D Dwyer	2 drugs/indications updated to reflect the date they move into routine commissioning
1.72	28-Mar-18	D Thomson; D Dwyer	1 drug/indication updated to reflect the date it moves into routine commissioning; 1 drug/indication moved into routine commissioning; 1 drug/indication with updated treatment criteria
1.73	03-Apr-18	P Clark; D Thomson; D Dwyer	1 drug/indication removed
1.74	09-Apr-18	P Clark; D Thomson; D Dwyer	1 drug/ indication for routine commissioning which will receive interim CDF funding
1.75	11-Apr-18	D Thomson; D Dwyer	1 drug/indication updated to reflect the date it moves into routine commissioning
1.76	19-Apr-18	P Clark; D Thomson; D Dwyer	1 drug/indication with updated treatment criteria
1.77	24-Apr-18	D Thomson; D Dwyer	2 drugs/indications moved into routine commissioning
1.78	25-Apr-18	D Thomson; D Dwyer	2 drugs/indications moved into routine commissioning
1.79	27-Apr-18	P Clark; D Thomson; D Dwyer	1 drug/indication added to the CDF
1.80	01-May-18	D Thomson; D Dwyer	1 drug/indication moved into routine commissioning
1.81	04-May-18	P Clark; D Thomson; D Dwyer	5 drugs/indications which will receive interim CDF funding; 2 drugs/indications for routine commissioning
1.82	16-May-18	D Thomson; D Dwyer	1 drug/indication updated to reflect the date it moves into routine commissioning
1.83	17-May-18	P Clark; D Thomson; D Dwyer	1 drug/ indication for routine commissioning which will receive interim CDF funding
1.84	25-May-18	P Clark; D Thomson; D Dwyer	1 drug/indication added to the CDF
1.85	01-Jun-18	P Clark; D Thomson; D Dwyer	1 drug/indication added to the CDF
1.86	05-Jun-18	D Thomson; D Dwyer	1 drug/indication moved into routine commissioning
1.87	13-Jun-18	P Clark; D Thomson; D Dwyer	8 drugs/indications updated to reflect the date they move into routine commissioning; 2 drugs/indications updated to note EMA recommendation; 1 drug/indication with updated treatment criteria
1.88	19-Jun-18	D Thomson; D Dwyer	2 drugs/indications moved into routine commissioning
1.89	26-Jun-18	D Thomson; D Dwyer	1 drug/indication moved into routine commissioning
1.90	28-Jun-18	P Clark; D Thomson; D Dwyer	1 drug/ indication for routine commissioning which will receive interim CDF funding; 1 drug/indication with updated treatment criteria
1.91	05-Jul-18	D Thomson; D Dwyer	2 drugs/indications with updated treatment criteria
1.92	10-Jul-18	D Thomson; D Dwyer	1 drug/indication moved into routine commissioning
1.93	12-Jul-18	P Clark; D Thomson; D Dwyer	2 drugs/ indications for routine commissioning which will receive interim CDF funding; 3 drugs/indications moved into routine commissioning; 1 drug/indication with updated treatment criteria
1.94	13-Jul-18	D Thomson; D Dwyer	1 drug/indication moved into routine commissioning;
1.95	20-Jul-18	P Clark; D Thomson; D Dwyer	1 drug/indication updated to reflect the date it moves into routine commissioning; 1 drug/ indication for routine commissioning which will receive interim CDF funding; 1 drug/indication with updated treatment criteria
1.96	25-Jul-18	P Clark; D Thomson; B Groves	1 drug in 2 indications entering a CDF managed access period
1.97	03-Aug-18	D Thomson; D Dwyer	1 drug/indication with updated treatment criteria
1.98	09-Aug-18	P Clark; D Thomson; D Dwyer	2 drugs/indications for routine commissioning which will receive interim CDF funding; 2 drugs/indications with updated treatment criteria; 3 drugs/indications updated to reflect the date they move into routine commissioning
1.99	14-Aug-18	B Groves; P Clark; D Thomson	1 drug/indication moved into routine commissioning; 1 drug/indication moved back to the CDF list
1.100	24-Aug-18	P Clark; D Thomson; D Dwyer	1 drug/indication for routine commissioning which will receive interim CDF funding; 3 drugs/indications with updated treatment criteria; 2 drugs/indications updated to reflect the date they move into routine commissioning

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1.101	31-Aug-18	P Clark; D Thomson; D Dwyer	2 drugs/indications with updated treatment criteria; 1 drug/indication updated to reflect the date it moves into routine commissioning
1.102	07-Sep-18	P Clark; D Thomson; D Dwyer	1 drug/indication moved into routine commissioning; 1 drugs/indications with updated treatment criteria
1.103	11-Sep-18	D Thomson; D Dwyer	7 drugs/indications moved into routine commissioning
1.104	17-Sep-18	P Clark; D Thomson; D Dwyer	1 drug/indication for routine commissioning which will receive interim CDF funding
1.105	05-Oct-18	P Clark; D Thomson; D Dwyer	2 drugs/indications for routine commissioning which will receive interim CDF funding; 2 drugs/indications updated to reflect the date it moves into routine commissioning; 1 drug/indication with an updated form code; 2 drugs/ indications with updated treatment criteria
1.106	16-Oct-18	P Clark; D Thomson; D Dwyer	1 drug/indication moved into routine commissioning; 18 drugs/indications with updated treatment criteria
1.107	06-Nov-18	P Clark; D Thomson; D Dwyer	2 drugs/indications moved into routine commissioning
1.108	08-Nov-18	P Clark; D Thomson; D Dwyer	2 drugs/ Indications for routine commissioning which will receive interim CDF funding
1.109	20-Nov-18	P Clark; D Thomson; D Dwyer	2 drugs/indication added to the CDF; 2 drugs/indications updated to reflect the date it moves into routine commissioning; 2 drugs/indications moved into routine commissioning
1.110	22-Nov-18	P Clark; D Thomson; D Dwyer	1 drug/indication added to the CDF
1.111	27-Nov-18	D Thomson; D Dwyer	1 drug/indication moved into routine commissioning
1.112	30-Nov-18	P Clark; D Thomson; D Dwyer	1 drug/indication added to the CDF
1.113	07-Dec-18	P Clark; D Thomson; D Dwyer	1 drug/indication added to the CDF; 1 drug/indication recommended for routine commissioning which will be available via a free of charge compassionate access scheme until 90 days after the date NICE publishes final guidance; 1 drug/indication updated to reflect the date it will be delisted; 1 drug/indication with updated treatment criteria
1.114	12-Dec-18	P Clark; D Thomson; D Dwyer	1 drug/indication with updated treatment criteria
1.115	17-Dec-18	P Clark; D Thomson; D Dwyer	3 drugs/indications with updated treatment criteria; 1 drug/indication updated to reflect the date it will be delisted
1.116	19-Dec-18	P Clark; D Thomson; D Dwyer	2 drugs/indications moved into routine commissioning; 2 drugs/indications with updated treatment criteria; 2 drugs/indications updated to reflect the date it moves into routine commissioning
1.117	21-Dec-18	P Clark; D Thomson; D Dwyer	3 drugs/indications with updated treatment criteria
1.118	31-Dec-18	P Clark; B Groves	8 drugs/indications updated; 1 drug/indication moved to routine commissioning
1.119	15-Jan-19	P Clark; D Dwyer	1 drug/indication moved to routine commissioning; 1 drug/indication removed from the CDF list; 1 drug/indication updated to reflect the date it moves into routine commissioning
1.120	17-Jan-19	P Clark; D Dwyer	1 drug/ indication for routine commissioning which will receive interim CDF funding; 2 drugs/indications with updated treatment criteria
1.121	18-Jan-19	P Clark; S Williamson; D Dwyer	2 drugs/ indications for routine commissioning which will receive interim CDF funding; 4 drugs/indications with updated treatment criteria
1.122	23-Jan-19	P Clark; S Williamson; D Dwyer	2 drugs/indications with updated treatment criteria
1.123	24-Jan-19	P Clark; S Williamson; D Dwyer	1 drug/indication with updated treatment criteria
1.124	25-Jan-19	P Clark; S Williamson; D Dwyer	2 drugs/indications suspended from CDF funding for new patients
1.125	01-Feb-19	P Clark; S Williamson; D Dwyer	1 drug/indication added to the CDF
1.126	01-Feb-19	P Clark; S Williamson; D Dwyer	2 drug/indication added to list B
1.127	15-Feb-19	P Clark; S Williamson; D Dwyer	1 drug/indication removed from the CDF; 2 drugs/indications moved to routine commissioning; 3 drugs/indications for routine commissioning which will receive CDF interim funding; 6 drugs/indications with updated treatment criteria
1.128	12-Mar-19	P Clark; S Williamson; D Dwyer	1 drug/indication added to the CDF; 3 drugs/indications updated to reflect the date it moves into routine commissioning
1.129	21-Mar-19	P Clark; S Williamson; D Dwyer	1 drug/ indication for routine commissioning which will receive interim CDF funding; 2 drugs/indications moved to rountine commissioning; 1 drug/indication with updated treatment criteria
1.130	28-Mar-19	P Clark; S Williamson; D Dwyer	1 drug/indication added to the CDF
1.131	02-Apr-19	P Clark; S Williamson; D Dwyer	1 drug/indication added to the CDF
1.132	05-Apr-19	P Clark; S Williamson; D Dwyer	1 drug/indication added to the CDF
1.133	09-Apr-19	P Clark; S Williamson; D Dwyer	1 drug/indication added to list 8; 1 drug/indication with updated treatment criteria
1.134	18-Apr-19	P Clark; S Williamson; D Dwyer	2 drugs/indications with updated treatment criteria; 3 drugs/indications updated to reflect the date it moves into routine commissioning
1.135	02-May-19	P Clark; S Williamson; D Dwyer	2 drugs/indications for routine commissioning which will receive interim CDF funding; 1 drug/indication updated to reflect the date it moves into routine commissioning
1.136	17-May-19	P Clark; S Williamson; D Dwyer	2 drugs/indications for routine commissioning which will receive interim CDF fundication with updated treatment criteria; 2 drugs/indication with new Blueteq forms created
1.137	28-May-19	P Clark; S Williamson; D Dwyer	a digg/ midications moved into routine commissioning
1.138	18-Jun-19	P Clark; S Williamson; D Dwyer	3 drugs/indications moved into routine commissioning
1.139	19-Jun-19	P Clark; S Williamson; D Dwyer	2 drugs/indications for routine commissioning which will receive interim CDF funding; 9 drug/indication with updated treatment criteria
1.140	02-Jul-19	P Clark; S Williamson; D Dwyer	a diagnostic in the commendation to the CDF
1.141	02-Jul-19 05-Jul-19	P Clark; S Williamson; D Dwyer P Clark; S Williamson; D Dwyer	2 drug/microstorio for routine commissioning which will receive interim CDF funding; 1 drug/indication moved to routine commissioning
1.142	17-Jul-19	P Clark; S Williamson; D Dwyer	La log michaetion in obtained minimal meter members and an interest members of the control of th
1.142	23-Jul-19	P Clark; S Williamson; D Dwyer	2 drugs/indications moved into routine commissioning
1.143	26-Jul-19	P Clark; S Williamson; D Dwyer P Clark; S Williamson; D Dwyer	2 drugs/motions under dot to reflect the date it moves into routine commissioning: 1 drug/indication recommeded to the CDF
1.144	30-Jul-19	P Clark; S Williamson; D Dwyer P Clark; S Williamson; D Dwyer	2 diagy/motations updated to renect the date in moves into double commissioning. I diagy/motation recommended to the COP  I drug/motation updated to reflect the date supply became available
1.145			1. Diagramukation upuateu to renets: the usate supply use centre available.  3. drugs/indications with updated treatment criteria.
	02-Aug-19	P Clark; S Williamson; D Dwyer	
1.147 1.148	06-Aug-19	P Clark; S Williamson; D Dwyer	1 drug/ Indication for routine commissioning which will receive interim CDF funding 1 drug/ Indication added to the CDF
	08-Aug-19	P Clark; S Williamson; D Dwyer	1 drug/mication added to the CDF
1.149	03-Sep-19	P Clark; S Williamson; D Dwyer	A string minutation above to the Con

Version No.	Date published	Author(s)	Revision summary
1.150	24-Sep-19	P Clark; S Williamson; D Dwyer	2 drug/indication added to list B
1.151	03-Oct-19	P Clark; S Williamson; D Dwyer	1 drug/indication updated to reflect the date supply became available
1.152 1.153	11-Oct-19 22-Oct-19	P Clark; S Williamson; D Dwyer	2 drugs/indications added to the CDF; 2 drugs/indications with updated treatment criteria
1.154	12-Nov-19	P Clark; S Williamson; D Dwyer P Clark; S Williamson; D Dwyer	2 drug/Indication added to list 8  I drug/Indication added to list 8, 7 drugs/Indications with updated criteria; 1 drug/Indication with treatment criteria added to list 8
1.155	28-Nov-19	P Clark; S Williamson; D Dwyer	1 drugs/indications added to the CDF, 2 drugs/indications with updated treatment criteria
1.156	29-Nov-19	P Clark; S Williamson; D Dwyer	1 drugs/indications added to the CDF; 1 drug/ indication for routine commissioning which will receive interim CDF funding; 6 drugs/indications with updated treatment criteria
1.157	04-Dec-19	P Clark; S Williamson; D Dwyer	4 drugs/indications with updated treatment criteria
1.158	15-Jan-20	P Clark; S Williamson; D Dwyer	1 drug/Indication for routine commissioning which will receive interim CDF funding; 4 drugs/Indications with updated treatment criteria
1.160	27-Feb-20 09-Mar-20	P Clark; S Williamson; D Dwyer P Clark; S Williamson; D Dwyer	1 drug/Indication added to list 8; 1 drug/Indication for routine commissioning which will receive interim CDF funding 3 drugs/Indications with updated treatment criteria
1.161	03-Apr-20	P Clark; S Williamson; D Dwyer	3 drug/miciation added to the CDF; 12 drugs/indications with updated treatment criteria
1.162	17-Apr-20	P Clark; S Williamson; D Dwyer	1 drug/indication recommended for the CDF; 17 drug/indications added to list C; 1 drug/indication added to list B
1.163	07-May-20	P Clark; S Williamson; D Dwyer	1 drug/indication for routine commissioning which will receive interim CDF funding; 17 drug/indications added to list C
1.164 1.165	22-May-20	P Clark; S Williamson; D Dwyer	1 drug/indication for routine commissioning which will receive interim CDF funding; 1 drug/indications added to list C; 6 drugs/indications with updated treatment criteria
1.166	27-May-20 13-Jul-20	P Clark; S Williamson; D Dwyer P Clark; S Williamson; D Dwyer	1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding; 2 drug/indications with updated treatment criteria; 1 drug/indication added to list B; 1 drug/indication with CDF exit date added
1.167	31-Jul-20	P Clark; S Williamson; D Dwyer	1 drug/indication added to the CDF; 1 drug/indication for routine commissioning which will receive interim CDF drug/indication added to list B; 1 drug/indication removed from list C
1.168	20-Aug-20	P Clark; S Williamson; D Dwyer	1 drug/Indication for routine commissioning which will receive interim CDF funding: 3 drugs/indications with published treatment criteria after marketing authorisation; 2 drugs/indications added to list 8; 4 drugs/indications with date moving to routine commissioning updated
1.169	11-Sep-20	P Clark; S Williamson; D Dwyer	2 drugs/indications for routine commissioning which will receive interim CDF funding; 6 indications added to list C; 1 drug/indication removed from list C; 5 drugs/indications with updated treatment criteria
1.170	23-Oct-20	P Clark; S Williamson; D Dwyer	2 drugs/indications added to the CDF; 1 drugs/indications for routine commissioning which will receive interim CDF funding; 1 indications removed from list C; 2 drugs/indications with updated treatment criteria
1.171	12-Nov-20	P Clark; S Williamson; D Dwyer	3 drugs/indications for routine commissioning which will receive interim CDF funding; 1 drugs/indications added to the CDF; 4 drugs/indications added to list B
1.172	25-Nov-20	P Clark; S Williamson; D Dwyer	1 drug/indication for routine commissioning which will receive interim CDF funding; 1 drugs/indications removed from list C; 2 drugs/indications with date moving to routine commissioning updated
1.173	15-Dec-20	P Clark; S Williamson; D Dwyer	3 drugs/indications for routine commissioning which will receive interim CDF funding; 5 drugs/indications with updated treatment criteria
1.174	19-Jan-21	P Clark; S Williamson; D Dwyer	3 drugs/indications added to the CDF; 3 drugs/indications added to list 8; 5 drugs/indications with updated treatment criteria
1.175 1.176	27-Jan-21 18-Feb-21	P Clark; S Williamson; D Dwyer P Clark; S Williamson; D Dwyer	1 drug/indication for routine commissioning which will receive interim CDF funding; 2 drugs/indications with updated treatment criteria  13 drugs/indications with updated treatment criteria; 1 drug/indication with an updated form title; 1 drug/indication updated to reflect the date it leaves the CDF after terminated guidance
1.177	19-Mar-21	P Clark; S Williamson; D Dwyer	13 urgs/mulations with updated treatment criteria, 2 urgs/mulations with updated treatment criteria, 2 urgs/mulations with updated treatment criteria; 4 drugs/indications soft outside to a feet terminated guidance.  2 drugs/mulations for routine commissioning which will receive interim CDF indication recommended for the CDF; 1 drugs/indications with updated treatment criteria; 4 drugs/indications added to list S
1.178	29-Mar-21	P Clark; S Williamson; R Mishra	9 drugs/indications removed from list C
1.179	28-Apr-21	P Clark; S Williamson; D Dwyer	2 durgs/indications removed from the CDF; 1 drug/indication recommended for the CDF; 1 drug/indication for routine commissioning which will receive interim CDF funding; 2 drugs/indications added to list B; 6 drugs/indications with updated date moving to routine commissioning
1.180	17-May-21	P Clark; S Williamson; D Dwyer	1 drug/indication added to list D; 2 drugs/indications recommended for routine commissioning; 1 drug/indication removed from list C; 7 drugs/indications with updated treatment criteria
1.181	17-Jun-21	P Clark; S Williamson; D Dwyer	2 drugs/midications for outside commissioning which will receive interim CDF funding: 11 drugs/midication and to list B; 8 drugs/midications with updated reatment criteria; 1 drugs/midication removed from list C; 1 drugs/midication removed from the CDF
1.182	25-Jun-21	P Clark; S Williamson; D Dwyer	1 drug/Indication removed from list B; 5 drugs/indications with updated treatment criteria
1.183	01-Jul-21	P Clark; S Williamson; D Dwyer	4 drugs/indications removed from list C; 1 drug/indication added to list B
1.184	23-Jul-21	P Clark; S Williamson; D Dwyer	1 drugs/indications for routine commissioning which will receive interin CDF funding; 1 drug/indication added to list 8; 7 drugs/indications with updated treatment criteria; 1 drug/indication removed from list C
1.185 1.186	30-Jul-21 21-Aug-21	P Clark; S Williamson; D Dwyer P Clark; S Williamson; D Dwyer	1 drugs/indications for routine commissioning which will receive interim CDF funding; 1 drug/indication added to list B; 1 drug/indication removed from list C  1 drugs/indications for routine commissioning which will receive interim CDF funding; 1 drug/indication with updated treatment criteria
1.187	10-Sep-21	P Clark; S Williamson; D Dwyer	2 drugs/midications for routine commissioning which will receive interim CD triunging 2 drugs/midications for routine commissioning which will receive interim CD triunging 2 drugs/midications for routine commissioning which will receive interim CD fruinging 2 drugs/midication with updated treatment criteria
1.188	17-Sep-21	P Clark; S Williamson; D Dwyer	1 drugs/indications for routine commissioning which will receive interim CDF funding; 1 drug/indication added to list B
1.189	21-Sep-21	P Clark; S Williamson; D Dwyer	1 drugs/Indications for routine commissioning which will receive interim CDF funding; 1 drug/indication added to list B; 4 drugs/indications with updated treatment criteria
1.190	24-Sep-21	P Clark; S Williamson; D Dwyer	1 drug/indication added to list B; 1 drug/indication with updated date moving to routine commissioning
1.191 1.192	01-Oct-21 08-Oct-21	P Clark; S Williamson; D Dwyer P Clark; S Williamson; D Dwyer	2 drugs/indications recommended for the CDF; 1 drug/indication with updated treatment criteria 2 drugs/indications added to list B; 1 drug/indication with an updated title
1.193	15-Oct-21	P Clark, S Williamson; D Dwyer	2 or ogymnications for routine commissioning which will receive interim CDF funding; 2 drugs/indications with updated treatment criteria
1.194	02-Nov-21	P Clark; S Williamson; D Dwyer	1 drug/indication added to list D; 1 drug/indication added to list B; 5 drugs/indications with updated date moving to routine commissioning
1.195	11-Nov-21	P Clark; S Williamson; D Dwyer	2 drugs/indications for routine commissioning which will receive interim CDF funding
1.196	17-Nov-21	P Clark; S Williamson; D Dwyer	1 drug/indication recommended for the CDF; 1 drug/indication with updated date moving to routine commissioning; 9 drugs/indications with updated treatment criteria
1.197 1.198	30-Nov-21 03-Dec-21	P Clark; S Williamson; D Dwyer P Clark; S Williamson; D Dwyer	2 drugs/indications recommended for the CDF; 2 drugs/indications with updated treatment criteria 5 drugs/indications with updated treatment criteria
1.198	16-Dec-21	P Clark; S Williamson; D Dwyer P Clark; S Williamson; D Dwyer	3 drugs/molcations with updated treatment criteria   drugs/molcations for routine commissioning which will receive interim CDF funding; 1 drug/indication with updated treatment criteria; 1 drug/indication added to list B; 1 drug/indication with updated date moving to routine commissioning
1.200	22-Dec-21	P Clark; S Williamson; D Dwyer	1 drug/indication for routine commissioning which will receive interim CDF funding; 8 drugs/indications with updated treatment criteria; 1 drug/indication added to list 8
1.201	21-Jan-22	P Clark; S Williamson; D Dwyer	1 drug/indication for routine commissioning which will receive interim CDF funding; 2 drugs/indications added to list B
1.202	26-Jan-22	P Clark; S Williamson; D Dwyer	3 drugs/indications added to list B
1.203 1.204	02-Feb-22 08-Feb-22	P Clark; S Williamson; D Dwyer P Clark; S Williamson; D Dwyer	1 drug/Indication added to list D, 3 drugs/Indications with updated date moving to routine commissioning 1 drug/Indication recommended for the CDF; 1 drug/Indication removed from list C
1.204	25-Feb-22	P Clark; S Williamson; D Dwyer P Clark; S Williamson; D Dwyer	1 oragination recommended for the CP; 1 drug/indication removes from ist 8  I drug/indication recommended for the CP; 1 drug/indication added to list 8
1.206	03-Mar-22	P Clark; S Williamson; D Dwyer	2 organisation recommended for the CDF; 2 drugs/indications added to list B
1.207	24-Mar-22	P Clark; S Williamson; D Dwyer	1 drug/indication recommended for the CDF; 2 drugs/indications added to list B: 10 drugs/indications with updated treatment criteria
1.208	01-Apr-22	P Clark; S Williamson; D Dwyer	7 drugs/indications removed from list C: 6 drugs/indications with updated treatment criteria
1.209 1.210	07-Apr-22	P Clark; S Williamson; D Dwyer	1 drug/Indication for routine commissioning which will receive interim CDF funding: 3 drugs/indications with updated treatment criteria    Advantable interior commissioning which will receive interim CDF funding: 3 drugs/indications with updated treatment criteria
1.210	14-Apr-22 05-May-22	P Clark; S Williamson; Z Niwaz P Clark; S Williamson; D Dwyer	2 drugs/indications for routine commissioning which will receive interim CDF funding; 9 drugs/indications with updated treatment criteria  1 drug/indication added to list D; 3 drugs/indications for routine commissioning which will receive interim CDF funding; 6 drugs/indications with updated treatment criteria
1.212	17-May-22	P Clark; S Williamson; D Dwyer	1 drug/microion added to 181, 3 drugs/microions with updated treatment criteria, 10 drugs/microion sudre under the criteria of
1.213	25-May-22	P Clark; S Williamson; Z Niwaz	2 drugs/indications for routine commissioning which will receive interim CDF funding; 2 drugs/indications added to list B; 1 drug/indication with updated treatment criteria
1.214	06-Jun-22	P Clark; S Williamson; Z Niwaz	6 drugs/indications with updated treatment criteria
1.215	17-Jun-22	P Clark; S Williamson; D Dwyer	1 drug/indication for routine commissioning which will receive interim CDF funding; 1 drug/indication moved into routine commissioning; 1 drug/indication removed from the CDF; 2 drugs/indications with updated dreatment criteria; 2 drugs/indications with updated date moving to routine commissioning
1.216	23-Jun-22	P Clark; S Williamson; Z Niwaz	1 drug/indication with updated date moving to routine commissioning; 3 drugs/indications moved into routine commissioning; 10 drugs/indications with updated treatment criteria
1.217	29-Jun-22	P Clark; S Williamson; Z Niwaz	1 drug/indication for routine commissioning which will receive interim CDF funding; 2 drugs/indications with updated date moving to routine commissioning; 1 drug/indication with updated treatment criteria
1.218	30-Jun-22 07-Jul-22	P Clark; S Williamson; Z Niwaz	1 drug/Indication for routine commissioning which will receive interim CDF funding
1.219	07-Jul-22 14-Jul-22	P Clark; S Williamson; Z Niwaz P Clark; S Williamson; Z Niwaz	1 drug/indication for routine commissioning which will receive interim CDF funding 3 drugs/indications for routine commissioning which will receive interim CDF funding; 1 drug/indication moved into routine commissioning; 3 drugs/indications with updated indication and treatment criteria
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Version No.	Date published	Author(s)	Revision summary
1.221	18-Jul-22	P Clark; S Williamson; Z Niwaz	1 drug/indication updated to reflect the date supply became available and treatment criteria added; 1 drug/indication with updated treatment criteria
1.222	20-Jul-22	P Clark; S Williamson; Z Niwaz	4 drugs/indications moved into routine commissioning; 1 drug/indication with updated date moving to routine commissioning; 2 drugs/indications with updated treatment criteria
1.223	26-Jul-22	P Clark; S Williamson; Z Niwaz	2 drugs/indications moved into routine commissioning
1.224	03-Aug-22	P Clark; S Williamson; Z Niwaz	1 drug/indication with updated date moving to routine commissioning
1.225	10-Aug-22	P Clark; S Williamson; Z Niwaz	2 drugs/indications with updated date moving to routine commissioning; 1 drug/indication with updated treatment criteria: changes made to section C and front page
1.226	18-Aug-22	P Clark; S Williamson; Z Niwaz	1 drug/indication for routine commissioning which will receive interim CDF funding; 1 drug/indication with updated date moving to routine commissioning; 1 drug/indication with updated treatment criteria
1.227	23-Aug-22 02-Sep-22	P Clark; S Williamson; Z Niwaz P Clark; S Williamson; Z Niwaz	1 drug/indication recommended for the CDF, removed from list D, with updated treatment criteria  1 drug/indication for routine commissioning which will receive interient CDF funding; 1 drug/indication moved into routine commissioning which will receive interient CDF funding; 1 drug/indication moved into routine commissioning which will receive interient CDF funding; 1 drug/indication moved into routine commissioning will receive the received the received funding to routine commissioning will receive the received funding the rece
1.228	07-Sep-22	P Clark; S Williamson; Z Niwaz	1 aroug/indication updated to reflect availability  1 droug/indication updated to reflect availability
1.230	16-Sep-22	P Clark; S Williamson; Z Niwaz	2 organisation opposed to tence designation and the service of the
1.231	23-Sep-22	P Clark; S Williamson; D Dwyer	1 drug/indication for routine commissioning which will receive interim CDF funding; 1 drug/indication with updated treatment criteria; 1 drug/indication moved into routine commissioning
1.232	07-Oct-22	P Clark; S Williamson; Z Niwaz	2 drugs/indications moved into routine commissioning; 1 drug/indication with updated date moving to routine commissioning; 1 drug/indication with updated treatment criteria
1.233	11-Oct-22	P Clark; S Williamson; Z Niwaz	1 drug/indication for routine commissioning which will receive interim CDF funding
1.234	13-Oct-22	P Clark; S Williamson; Z Niwaz	1 drug/indication updated to reflect the date supply became available
1.235	19-Oct-22	P Clark; S Williamson; Z Niwaz	2 drugs/indications moved into routine commissioning; 2 drugs/indications with updated date moving to routine commissioning; 3 drugs/indications removed from list C; 13 drugs/indications assigned with Blueteq Form references
1.236	26-Oct-22	P Clark; S Williamson; Z Niwaz	2 drugs/indications with updated date moving to routine commissioning
1.237	08-Nov-22	P Clark; S Williamson; Z Niwaz	1 drug/indication for routine commissioning which will receive interim CDF funding; 2 drugs/indications moved into routine commissioning
1.238	10-Nov-22	P Clark; S Williamson; Z Niwaz	2 drugs/indications for routine commissioning which will receive interim CDF funding; 2 drugs/indications with updated treatment criteria; 1 drug/indication with updated indication and treatment criteria
1.239	16-Nov-22	P Clark; S Williamson; Z Niwaz	1 drug/indication recommended for the CDF, removed from list D, with updated treatment criteria; 1 drug/indication moved into routine commissioning
1.240	24-Nov-22	P Clark; S Williamson; Z Niwaz	1 drug/indication for routine commissioning which will receive interim CDF funding
1.241	25-Nov-22	P Clark; S Williamson; Z Niwaz	1 drug/indication added to list D
1.242	14-Dec-22	P Clark; S Williamson; Z Niwaz	3 drugs/indications with updated date moving to routine commissioning
1.243	20-Dec-22	P Clark; S Williamson; Z Niwaz	1 drug/indication recommended for the CDF; 1 drug/indication with updated indication and treatment criteria
1.244	22-Dec-22	P Clark; S Williamson; Z Niwaz	1 drug/indication for routine commissioning which will receive interim CDF funding; 1 drug/indication assigned with a Blueteq Form reference; 1 drug/indication with updated indication; 2 drugs/indications with updated treatment criteria
1.245 1.246	04-Jan-23 12-Jan-23	P Clark; S Williamson; Z Niwaz P Clark; S Williamson; Z Niwaz	1 drug/indication with updated date moving to routine commissioning 2 drugs/indication with updated date moving to routine commissioning 3 drugs/indications with updated date moving to routine commissioning
1.246			
1.248	18-Jan-23 25-Jan-23	P Clark; S Williamson; Z Niwaz P Clark; S Williamson; Z Niwaz	2 drugs/indications moved into routine commissioning; 1 drug/indication for routine commissioning which will receive interim CDF funding; 1 drug/indication with updated treatment criteria  1 drug/indication for routine commissioning which will receive interim CDF funding; 2 drugs/indication with updated treatment criteria; 1 drug/indication with updated Blueteq Form reference
1.249	26-Jan-23	P Clark; S Williamson; Z Niwaz	La trag mutation for routine commissioning which will receive interim CDF funding; 1 drug/indication with updated treatment criteria, 1 drug/indication for routine commissioning which will receive interim CDF funding; 1 drug/indication with updated treatment criteria.
1.250	09-Feb-23	P Clark; S Williamson; Z Niwaz	a drug/indication with updated CDF managed access status, 2 drugs/indications with updated access status, 2 drugs/indications
1.251	22-Feb-23	P Clark; S Williamson; Z Niwaz	1 drug/indication for routine commissioning which will receive interim CDF funding; 1 drug/indication with updated CDF managed access status; 1 drug/indication with updated date moving to routine commissioning; 1 drug/indication with updated treatment criteria
1.252	01-Mar-23	P Clark; S Williamson; Z Niwaz	2 drugs/indications with updated date moving to routine commissioning; 2 drugs/indications with updated treatment criteria
1.253	09-Mar-23	P Clark; S Williamson; Z Niwaz	2 drugs/indications added to routine commissioning; 20 drugs/indications with updated treatment criteria
1.254	14-Mar-23	P Clark; S Williamson; Z Niwaz	3 drugs/indications moved into routine commissioning; 6 drugs/indications with updated treatment criteria
1.255	22-Mar-23	P Clark; S Williamson; Z Niwaz	1 drug/indication with updated date moving to routine commissioning
1.256	29-Mar-23	P Clark; S Williamson; Z Niwaz	1 drug/indication recommended for the CDF
1.257	31-Mar-23	P Clark; S Williamson; Z Niwaz	4 drugs/indications removed from list C; 2 drugs/indications with updated treatment criteria
1.258	06-Apr-23	P Clark; S Williamson; Z Niwaz	2 drugs/indications for routine commissioning which will receive interim CDF funding; 1 drug/indication moved into routine commissioning
1.259	11-Apr-23	P Clark; S Williamson; Z Niwaz	2 drugs/indications moved into routine commissioning; 2 drugs/indications (4 forms) with updated treatment criteria
1.260	21-Apr-23	P Clark; S Williamson; Z Niwaz	1 drug/indication for routine commissioning which will receive interim CDF funding; 1 drug/indication (2 forms) with updated treatment criteria
1.261	24-Apr-23	P Clark; S Williamson; Z Niwaz	1 drug/indication for routine commissioning which will receive interim CDF funding; 1 drug/indication with updated indication and treatment criteria
1.262	27-Apr-23	P Clark; S Williamson; Z Niwaz	2 drugs/indications recommended for the CDF; 1 drug/indication (2 forms) with updated drug name and treatment criteria
1.263	04-May-23	P Clark; S Williamson; Z Niwaz	1 drug/indication with updated Blueteq form reference; 6 drugs/indications with updated drug column; 6 drugs/indications with updated treatment criteria
1.264	11-May-23	P Clark; S Williamson; J Hill	1 drugs/indication for routine commissioning which will receive interim CDF funding, removed from list C; 2 drugs/indications moved into routine commissioning, with updated treatment criteria; 2 drugs/indications (4 forms) with updated date moving to routine commissioning
1.265	18-May-23	P Clark; S Williamson; J Hill	2 drugs/indications for routine commissioning which will receive interim CDF funding; 1 drug/indication with updated treatment criteria
1.266	02-Jun-23	P Clark; R Nijjar; J Hill	3 drugs/indications moved into routine commissioning; 1 drug/indication with updated date moving to routine commissioning; 2 drugs/indications with updated treatment criteria; 2 drugs/indications with updated Blueteq form reference; 1 drug/indication with updated drug column
1.267	08-Jun-23	R Nijjar; J Hill	1 drug/indication for routine commissioning which will receive interim CDF funding; 8 drugs/indications with updated Blueteq form reference
1.268	14-Jun-23	P Clark; S Williamson; J Hill	1 drug/indication with updated date moving to routine commissioning
1.269	22-Jun-23	P Clark; S Williamson; Z Niwaz	1 drug/indication recommended for the CDF; 1 drug/indication moved into routine commissioning; 2 drugs/indications with updated date moving to routine commissioning
1.270	31-Jul-23	P Clark; S Williamson; J Hill	2 drugs/indications with updated treatment criteria
1.271	08-Aug-23	P Clark; S Williamson; J Hill	2 drugs/indications (4 forms) moved into routine commissioning; 1 drug/indication with updated treatment criteria; 1 drug/indication with updated TA number, Date of final NICE guidance, Date baseline funding started
1.272	17-Aug-23	P Clark; S Williamson; J Hill	1 drug/indication (5 forms) for routine commissioning which will receive interim CDF funding; 1 drug/indication removed from list C
1.273	24-Aug-23	P Clark; S Williamson; J Hill	2 drugs/indications with updated treatment criteria
1.274	07-Sep-23	P Clark; J Hill	1 drug/indication for routine commissioning which will receive interim CDF funding; 2 drugs/indications moved into routine commissioning; 2 drugs/indications with updated Previous CDF drug/ indication column
1.275	12-Sep-23	P Clark; J Hill	1 drugs/indications moved into routine commissioning
1.276 1.277	14-Sep-23 22-Sep-23	P Clark; J Hill P Clark; J Hill	1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commissioning which will receive interim CDF funding  1 drug/indication for routine commission for routine commiss
1.277	22-Sep-23 19-Oct-23	P Clark; J Hill	1 drug/indication for routine commissioning which will receive interim CDF funding; 2 drug/indications moved into routine commissioning; 11 drugs/indications with updated treatment criteria; 5 drugs/indications with updated date moving to routine commissioning to routine commissioning which will receive interim CDF funding; 1 drug/indication with updated date moving to routine commissioning; 9 drugs/indications with updated treatment criteria; 1 drug/indication with updated 'Expected date moving to routine commissioning; 9 drugs/indications with updated treatment criteria; 1 drug/indication with updated 'Expected date moving to routine commissioning which will receive interim CDF funding; 1 drug/indication with updated date moving to routine commissioning; 9 drugs/indications with updated date moving to routine commissioning which will receive interim CDF funding; 1 drug/indication with updated date moving to routine commissioning which will receive interim CDF funding; 1 drug/indication with updated treatment criteria; 1 drug/indication with updated treatment criteria; 1 drug/indication with updated treatment criteria; 1 drug/indication with updated treatment criteria; 1 drug/indication with updated treatment criteria; 1 drug/indication with updated treatment criteria; 1 drug/indication with updated treatment criteria; 1 drug/indication with updated treatment criteria; 1 drug/indication with updated treatment criteria; 1 drug/indication with updated treatment criteria; 1 drug/indication with updated treatment criteria; 1 drug/indication with updated treatment criteria; 1 drug/indication with updated treatment criteria; 1 drug/indication with updated treatment criteria; 1 drug/indication with updated treatment criteria; 1 drug/indication with updated treatment criteria; 2 drug/indication with updated treatment criteria; 2 drug/indication with updated treatment criteria; 3 drug/indication with updated treatment criteria; 3 drug/indication with updated treatment criteria; 3 drug/indication with updated treatment
			Entry into Baseline Commissioning' status
1.279	01-Nov-23	P Clark; J Hill	1 drug/indication updated to reflect the date supply became available and treatment criteria added; 1 drug/indication with updated date moving to routine commissioning  1 drug/indication for poulton or promptication updated in the promptication of the promptication updated in the promptication of the promptication updated in the prom
1.280 1.281	17-Nov-23 23-Nov-23	P Clark; J Hill P Clark; J Hill	1 drug/indication for routine commissioning which will receive interim CDF funding with updated treatment in the property of t
1.281	23-Nov-23 30-Nov-23	P Clark; J Hill	1 drug/indication for routine commissioning which will receive interin the routine form with updated date moving to routine commissioning.  1 drug/indication commission with updated in the routine commission with updated date moving to routine commissioning.  1 drug/indication commission with CVIS-1 drug/indication addition. It is drug in the routine commission with updated date moving to routine commissioning.
1.283	08-Dec-23	P Clark; J Hill	1 drug/indication removed from the CDF; 1 drug/indication added to list 8; 1 drug/indication removed from the CDF; 1 drug/indication added to list 8; 1 drug/indication removed from list CDF; 1 drug/indication for routine commissioning which will receive interier IDCF fundications added to list 8; 1 drug/indication with updated treatment criteria
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1.307 1.308 1.309				2 wigginandarina marcu mo roume commissioning, a wiginandarina attit updated uste mornig to roume commissioning (z. minis)
1387 1349/24 Palls (Palls and Control 1997) 1349/24 Palls	1.306	10-May-24	P Clark; J Richardson; J Hill	1 drug/indication for routine commissioning which will receive interim CDF funding; 1 drug/indication added to list B; 2 drugs/indications moved into routine commissionine: 2 drugs/indications with updated treatment criteria: 1 drug/indication with updated date moving to routine commissioning
1388 2 1892-3 P. Chuk J. Rebuthors (1981 ) Engly-inductions moved into endour commissioning. 3 singly-induction from the special position of commissioning of the commissioning o	1.307	17-May-24	P Clark; J Richardson; J Hill	
1310 07-30-22 P. CRA. P. Richardon, 180 1 degly-closure moved in continuous process commissioners. 2 supplemental models and transmitter critical	1.308		P Clark; J Richardson; J Hill	
1311 13-0-20 F Chris, Ribandon, 1981 1, Segündanic spatial to risk the step by locate analysis of testing to risk of the step by locate analysis of testing to risk of the step by locate analysis of testing to risk of the step by locate analysis of testing to risk of the step by locate analysis of testing to risk of the step by locate analysis of the step by locate analy				
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1.156 Sep. 32 P. Char, 1964 control, 1961 In supplication for restinct commissioning with will receive interior CF funding, 3 design/fedications with updated restinance criterion.  1.157 On Aug. 24 P. Char, 1964 control, 1961 In September 1964 of the Character		16-Jul-24		
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1.320 23-Aug 24 P Clurk; Richardson; Hill of uniffication for tronscore commissioning which will receive interim CDF funding. 2 drugs/indications with updated indication column.  1.321 05-5p-24 P Clurk; Richardson; 1 Hill of uniffication for tronscore commissioning with will receive interim CDF funding. 2 drugs/indications with updated date moving to routine commissioning. 2 drugs/indications with updated date moving to routine commissioning. 2 drugs/indications with updated date moving to routine commissioning. 2 drugs/indications with updated date moving to routine commissioning. 2 drugs/indications with updated indication column. 4 drugs/indication routine date moving to routine commissioning. 2 drugs/indications with updated indication column. 4 drugs/indication indication individual of treatment criteria.  1.322 05-5p-24 P Clurk; Richardson; Hill of uniffication for tronscore commissioning. 1 drugs/indication with updated date moving to routine commissioning. 3 drugs/indications with updated indication column. 4 drugs/indications with up				
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1.353   07-Mar-25   P Clark; J Richardson; J Hill   1 drug/indication (2 forms) added to list b; 2 drugs/indications with updated treatment criteria	1.353	07-Mar-25	P Clark; J Richardson; J Hill	Long/monation with realisms criteria above, a timp group monation with updated treatment criterion   drug/molation (2 forms) added to list b; 2 drugs/indications with updated treatment criteria   drug/molation (2 forms) added to list b; 2 drugs/indications with updated treatment criteria
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Version No.	Date published	Author(s)	Revision summary
1.356	26-Mar-25	P Clark; J Richardson; J Hill	1 drug/indication for routine commissioning which will receive interim CDF funding; 2 drug/indications (3 forms) with updated date moving to routine commissioning
1.357	02-Apr-25	P Clark; J Richardson; J Hill	1 drug/indication for routine commissioning which will receive interim CDF funding; 2 drugs/indications with updated treatment criteria
1.358	10-Apr-25	P Clark; J Richardson; J Hill	2 drugs/indications for routine commissioning which will receive interim CDF funding; 2 drugs/indications with updated treatment criterion
1.359	11-Apr-25	P Clark; J Richardson; J Hill	1 drug/indication for routine commissioning which will receive interim CDF funding
1.360	25-Apr-25	P Clark; J Richardson; J Hill	2 drugs/Indications for routine commissioning which will receive interim CDF funding; 2 drugs/indications moved into routine commissiong; 2 drugs/indications with updated treatment criteria
1.361	02-May-25	P Clark; J Richardson; J Hill	8 drugs/Indications with updated treatment criteria
1.362	09-May-25	P Clark; J Richardson; J Hill	2 drugs/Indications moved into routine commissiong; 2 drug/Indications with updated date moving to routine commissioning
1.363	16-May-25	P Clark; J Richardson; J Hill	2 drugs/indications (4 forms) moved into routine commissiong; 5 drugs/indications with updated treatment criteria; 1 drug/indication with updated date moving to routine commissioning
1.364	23-May-25	P Clark; J Richardson; J Hill	1 drug/indication moved into routine commissiong; 6 drugs/indications with updated treatment criteria; 1 drug/indication with updated TA column
1.365	06-Jun-25	P Clark; J Richardson; J Hill	1 drug/indication moved into routine commissiong; 8 drugs/indications with updated treatment criteria; 1 drug/indication with updated date moving to routine commissioning
1.366	12-Jun-25	P Clark; J Richardson; J Hill	1 drug/indication for routine commissioning which will receive interim CDF funding
1.367	27-Jun-25	P Clark; J Richardson; J Hill	1 drug/indication for routine commissioning which will receive interim CDF funding; 1 drug/indication for routine commissioning which moved directly into section B; 2 drugs/indications moved into routine commissioning; 11 drugs/indications with updated treatment criteria; 1 drug/indication with updated date moving to routine commissioning
1.368	03-Jul-25	P Clark; J Richardson; J Hill	1 drug/indication for routine commissioning which will receive interim CDF funding; 2 drugs/indications with updated treatment criteria; 1 drug/indication with updated date moving to routine commissioning
1.369	25-Jul-25	J Richardson; J Hill	2 drugs/Indications (3 forms) moved into routine commissioning; 2 drugs/Indications with updated treatment criteria; 3 drugs/Indications with updated date moving to routine commissioning
1.370	29-Jul-25	J Richardson; J Hill	1 drug/indication for routine commissioning which will receive interim CDF funding; 1 drug/indication with updated treatment criteria
1.371	06-Aug-25	J Richardson; R Chauhan; J Hill	1 drug/indication (2 forms) for routine commissioning which will receive interim CDF funding; 2 drugs/indications moved into routine commissioning
1.372	21-Aug-25	J Richardson; R Chauhan; S Ahmed	1 drug/indication for routine commissioning which will receive interim CDF funding; 2 drugs/indications moved into routine commissiong; 1 drug/indication with updated treatment criterion
1.373	04-Sep-25	J Richardson; R Chauhan; J Hill	1 drug/indication for routine commissioning which will receive interim CDF funding; 2 drugs/indications moved into routine commission; 1 drug/indication (2 forms) with updated date moving to routine commissioning
1.374	16-Sep-25	J Richardson; J Hill	1 drug/indication with updated date moving to routine commissioning
1.375	07-Oct-25	J Richardson; J Hill	1 drug/indication for routine commissioning which will receive interim CDF funding; 1 drug/indication (4 forms) removed from CDF weblist, 1 drug/indication with updated treatment criterion; 4 drugs/indications with updated treatment criteria
1.376	24-Oct-25	J Richardson; J Hill	1 drug/indication for routine commissioning which will receive interim CDF funding; 6 drugs/indications with updated treatment criteria; 1 drug/indication moved into routine commission; 1 drug/indication with updated date moving to routine commissioning
1.377	13-Nov-25	S O'Brien; J Richardson; J Hill	1 drug/indication for routine commissioning which will receive interim CDF funding; 12 drugs/indications with updated treatment criteria; 1 drug/indication moved into routine commissiong; 1 drug/indication with updated date moving to routine commissioning
1.378	18-Nov-25	S O'Brien; J Richardson; Z Niwaz	1 drug/indication for routine commissioning which will receive interim CDF funding; 1 drug/indication moved into routine commissioning
1.379	26-Nov-25	S O'Brien; R Hudson; Z Niwaz	2 drugs/indications for routine commissioning which will receive interim CDF funding; 1 drug/indication (2 forms) moved into routine commissioning

#### Changes to recent versions

General or criteria	Summary of changes
changed	Summer of Changes
Changes to version 1.379	
DOS3	Recommended for routine commissioning, receiving CDF interim funding
OBE01a	Recommended for routine commissioning, receiving CDF interim funding
OBE01b	
PEMB32	Moved into routine commissioning - section 8 of list
PEMB33	
Changes to version 1.378	
TALQ1	Recommended for routine commissioning, receiving CDF interim funding
DUR6	Moved into routine commissioning - section B of list
Changes to version 1.377	
GLO2	Recommended for routine commissioning, receiving CDF interim funding
ATE9	Treatment criteria (#6 and 12) updated
BLI2	Treatment criteria (#5, 7 and 10) updated
BLI3	Treatment criterion (#10) updated
BLI4	Treatment criterion (#10) updated
BLI5	Treatment criterion (#11) updated
BLI6	Treatment criterion (#11) updated
LNV3	Treatment criteria (#4 and 8) updated
NIV8a	Treatment criterion (#4) updated
NIV18	Treatment criterion (#4) updated
PEMB2	Treatment criteria (#6, 8 and 12) updated
PEMB9a	Treatment criteria (#4 and 10) updated
SOR3	Treatment criteria (#4 and 8) updated
RIB3	Moved into routine commissioning - section 8 of list
DARO3	Date moving into routine commissioning updated
Changes to version 1.376	
DARO3	Recommended for routine commissioning, receiving CDF interim funding
ABI4	Treatment criteria (#6 and 10) updated
APA2	Treatment criteria (#8 and 10) updated
ELR1	Treatment criteria (#9 and 15) updated
ENZ3	Treatment criteria (#7 and 9) updated
NIV21	Treatment criteria (#9 and 13) updated
NIV24	Treatment criteria (#2, 9 and 11) updated
FRU1	Moved into routine commissioning - section 8 of list
LOR2	Date moving into routine commissioning updated
Changes to version 1.375	
LOR2	Recommended for routine commissioning, receiving CDF interim funding
NHSE Urgent Interim	Removed from CDF weblist
Commissioning Policy	
Proposition 2420	
ALE1	Drug column updated; Treatment criterion (#4) updated; Treatment criteria (#8 and 10) removed
BRI2	Drug column updated; Treatment criterion (#4) updated; Treatment criteria (#8, 9 and 11) removed
DUR7	Treatment criterion (#7) and date moving into routine commissioning updated
ISA2	Date moving into routine commissioning updated
Changes to version 1.374	
DUR7	Recommended for routine commissioning, receiving CDF interim funding
ENF1	Date moving into routine commissioning updated
Changes to version 1.373	
ISA2	Recommended for routine commissioning, receiving CDF interim funding
CAP1	Moved into routine commissioning - section B of list
DOS2	Moved into routine commissioning - section B of list
NIV24	Moved into routine commissioning - section B of list
PEMB32	Date moving into routine commissioning updated
PEMB33	
Changes to version 1.372	
ENF1	Recommended for routine commissioning, receiving CDF interim funding
ERD1	Moved into routine commissioning - section B of list
ZAN6	Moved into routine commissioning - section B of list
FRU1	Treatment criterion (#4) updated
Changes to version 1.371	
PEMB32	Recommended for routine commissioning, receiving CDF interim funding
PEMB33	
BRE15	Moved into routine commissioning - section B of list
OSI4	Moved into routine commissioning - section B of list
Changes to version 1.370	
DUR6	Recommended for routine commissioning, receiving CDF interim funding
ATE8	Treatment criteria (#2, 5 and 11) updated