

	pecification: Vincristine 50ml infusion bags
Name of product	Vincristine infusion. Aseptically prepared from licensed sterile starting materials
Concentration	Concentration range 0.02mg/ml to 0.1mg/ml in 50ml infusion bags in accordance
	the national dose banding tables.
	The Vincristine concentrate should be added to the bag without withdrawal of
	equivalent volume of Sodium Chloride 0.9% w/v Infusion unless withdrawal of so
	infusion fluid is required to accomodate the total dose.
Diluent	Sodium Chloride 0.9% w/v
Volume	50ml
Final container	Non-PVC e.g. polyolefin infusion bag with additive port cover.
	Ideally infusion bag design will incorporate self sealing giving port to minimise F
	risks associated with accidental spillage during administration.
Starting materials	Licensed Vincristine Injection
	Licensed Sodium Chloride 0.9% w/v Infusion bags
Labelling	Labelling must be compliant with the principles of labelling for safety (in particul
	comply with HSC 2008/001 Updated national guidance on the
	safe administration of intrathecal chemotherapy and NPSA/2008/RRR004 i.e. m
	labelled 'For Intravenous Use Only – Fatal If Given by Other Routes') and the Bl
	General specification on unlicensed medicines. Tall Man lettering must be used
	drug name.
Label sample	An example label is provided below stating the minimum requirements only (the format is not restrictive and suppliers can use their preferred layout):-
	vinCRIStine xmg in xxml Sodium Chloride 0.9% w/v
	FOR INTRAVENOUS USE ONLY
	FATALIF GIVEN BY OTHER ROUTES
	Infuse the entire contents of the bag
	Store in a Refrigerator at 2-8°C Protect From Light
	Expiry: dd/mm/yyyy BN: XXXXXXXXX Keep out of the reach and sight of children
	Manufacture r's details MS XXXXXX
	Caution Cytotoxic: Handle with care
Batch Number	All products will have a unique batch identification number
Latex status of	All materials and manufacturing processes will be latex free or clearly labelled in
- components	
- manufacturing process	
Stability	Stability studies should conform to the Standard Protocol for deriving and asses of stability of Aseptic preparations (small molecules) published by the NHS