Clinical Practice Summary

Guidance on consensus approaches to managing Palliative Care Symptoms

Lancashire and South Cumbria Consensus Guidance - August 2017
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Alternatively- The Supportive & Palliative Care Indicators Tool (SPICT™) is available via link http://www.spict.org.uk

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Every effort has been made to ensure the accuracy of this text. However, the editorial team do not accept responsibility or legal liability for any errors in the text, or for the misuse or misapplication of material in this work.

Clinical Practice Summary For Palliative Care Symptoms

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These practice summaries are a place to begin.

They cannot replace advice from experienced clinicians.

Fundamental to the practice of palliative and end of life care is the individualised care of the patient and those important to them. If symptoms fail to respond to usual measures, or you are concerned that the guidance here may not be appropriate to the clinical situation you are in, contact your local specialist palliative care service for advice.

IF IN DOUBT ASK.

Background

In 2012 Lancashire and South Cumbria Specialist Palliative Care group wrote prescribing guidelines around managing common symptoms in a palliative care setting. These were well received and in 2014 were updated. In 2016, Lancashire and South Cumbria joined with Mersey and Cheshire in a new Strategic Clinical Network based around the North West Coast. As a result, this new version was developed, based on the guidance produced by our neighbouring Northern Strategic Clinical Network’s Guidelines (2016) and a Mersey and Cheshire Clinical Practice Summary (2017).

We have worked hard to try and achieve consensus and base the practice summaries on the best available evidence. We hope that in doing this we can help to ensure a consistency of approach to managing common symptoms, particularly for those individuals who receive care in a number of different locations.

Whilst every care has been taken to ensure accuracy and clarity, prescribers and clinicians must make all their decisions based on a full clinical assessment and their assessment of the risks and benefits of any intervention. They must also take into account any local guidance where it exists. In some areas the first line injectable opioid is Diamorphine not Morphine, contact your local Specialist Palliative Care team if advice required.

The evidence-base for prescribing in palliative care is not extensive or robust, which means that some guidance is based on a consensus of expert opinion. Many medications are used beyond licence and at doses that differ from other areas of clinical practice. This makes it impossible to produce guidance that contains definitive statements about what to prescribe and when.

Key Expert Resources:


Useful websites

Advance Care Planning

Advance Care Planning—North West Coast initiative

Deciding Right—North East initiative around Advance Care Planning
www.nescn.nhs.uk/common-themes/deciding-right

Recommended summary plan for emergency care and treatment (ReSPECT) - National Resuscitation Council Guidelines around summary care plans about patient’s preferences for care
www.respectprocess.org.uk

Knowledge Hub around end of life care and medication
http://endoflifecareambitions.org.uk/
http://www.palliativedrugs.com/

NICE guidance

Care of the dying adult in last days of life (2015) www.nice.org.uk/guidance/ng31

End of life care for infants, children and young people with life-limiting conditions: planning and management (2016)
www.nice.org.uk/guidance/ng61

Palliative care for adults: strong opioids for pain relief (2016)
www.nice.org.uk/guidance/cg140

Introduction and Aide Memoire

These easy reference guidelines are based on the Merseyside and Cheshire Palliative Care Network Audit Group Guidelines, Northern England Strategic Clinical Network Palliative and End of Life Care Guidelines 2016 and the Lancashire and South Cumbria Palliative Care Prescribing Guidelines 2014. They support decision-making in symptom management and care co-ordination for people in the last weeks of their life. If there is any doubt regarding clinical decisions for individuals, help should be sought from local Specialist Palliative Care services.

Ambitions for Palliative and End of Life Care - supporting people in the last weeks of life

All approaches regarding palliative and end of life care should reflect Ambitions for Palliative and End of Life Care, a national framework for local action 2015-2020 and the 6 key principles

- Each person is seen as an individual and
- Receives fair access to care
- We maximise comfort & wellbeing
- Care is coordinated
- All staff are prepared to care
- Each community is prepared to help

Ensure that you have considered the following in communication with the person and those important to them:

Preferences and possibilities that could constitute an Advance Care Plan

Sensitive communication about care in the last days of life including decisions about Do Not Attempt Cardiopulmonary Resuscitation (DNACPR) Orders or “allow natural death” decisions. Record these decisions and share with key organisations including “out of hours” care providers via Electronic Palliative Care Coordination System (EPaCCS) in line with local policies.

Ensure that there is a plan for the management of complex interventions such as non–invasive ventilation or Implantable Cardioverter Defibrillator (ICD) is in place, so they can be safely withdrawn when it is appropriate to do so.

Ensure that all relevant Out of Hours services are made aware of any critical documentation e.g. using special note notification in community or in hospital settings, that clear treatment escalation plans are made

Anticipatory prescribing to relieve common symptoms in the last weeks of life should be considered in a timely manner and individualised to avoid delay in managing distressing symptoms (Care of dying adults in the last days of life, NICE guideline NG31)

One Chance to Get it Right - Care in the last days and hours of life

Recognise deterioration and consider if this is potentially reversible, e.g. infection, or if the person is likely to die from irreversible causes. Potentially reversible causes should be treated provided that this is in accordance with the person’s wishes or in their best interests.

If the person is likely to die from irreversible causes in the next hours or few days communicate this clearly and sensitively.

Involv the dying person and those important to them in day-to-day decisions about personal care and clinical treatments.

Avoid undertaking investigations that are unlikely to affect care in the last few days of life unless there is a clinical need to do so (NG31) e.g. curtailing renal monitoring in advanced heart failure.

Construct an individual plan of care, which includes food and drink, symptom control and psychological, social and spiritual support.

Deliver this plan of care sensitively and review frequently, especially if symptoms are not controlled, there is concern from family members or the person shows sign of improvement

(Hydration is not covered in these guidelines but guidance can be found in the NICE Guidance NG31)
The North West End of Life Care Model

Supporting the people of the North West to live well before dying with peace and dignity in the place of their choice

End of life care

- Is about the individual and those important to them
- Is about meeting the supportive and palliative care needs for all those with an advanced progressive incurable illness or frailty, to live as well as possible until they die.
- Support may be needed in the last years, months or days of life.

It should include:

- A person centred approach to care - involving the person, and those closest to them in all aspects of their care including the decision making process around treatment and care
- Open, honest and sensitiv communication with the patient and those important to them
- Care which is coordinated and delivered with kindness and compassion
- The needs of those identified as important to the person to be actively explored, respected and met as far as possible
- All discussions to follow guidance set within the Mental Capacity Act (MCA 2005)

Key recommended Training for health and care staff:

Communication skills
Holistic assessment to include: physical, psychological, spiritual and social care
Symptom control
Advance care planning
Caring for carers
Priorities for care of the dying person
Bereavement support
Mental Capacity Act

The model supports the assessment and planning process for patients from the diagnosis of a life limiting illness or those who may be frail. The model comprises 5 phases and the Good Practice Guide (overleaf) identifies key elements of practice within each phase to prompt the assessment process as relevant to each setting.
## End of Life Care Good Practice Guide

### LAST YEAR OF LIFE

**Year/s**

- Patient identified as deteriorating despite optimal therapeutic management of underlying medical condition(s)
- Clear, sensitive communication with patient and those identified as important to them
- Person and agreed others are involved in decisions about treatment and care as they want
- Needs of those identified as important are explored, respected and met as far as possible
- Patient included on Supportive Care Record / GP Gold Standards Framework register and their care reviewed regularly
- Request consent to share information and create EPaCCS record
- Holistic needs assessment: physical, psychological, spiritual & social
- Keyworker identified
- Identify when there is an opportunity to offer an Advance Care Planning discussion. PPC/ADRT/LPA
- Making a will
- DNACPR discussion if appropriate
- Benefits review of patient and carer including: grants/prescription exemption
- Provide information on Blue Badge (disabled parking) scheme
- Agree on-going monitoring and support to avert crisis
- Referral to other services e.g. Specialist Palliative Care
- OOH/NWAS updated including Advance Care Plan/DNACPR
- ICD discussion if applicable

### INCREASING DECLINE

**Months/Weeks**

- Medical review
- All reversible causes of deterioration explored
- Clear, sensitive communication with patient and those identified as important to them
- Person and agreed others are involved in decisions about treatment and care as they want
- Needs of those identified as important are explored, respected and met as far as possible
- Prioritised as appropriate at Gold Standards Framework meeting
- On-going District Nurse support
- Agree on-going monitoring and support to avert crisis
- Holistic needs assessment
- Ongoing communication with Keyworker
- Review or offer advance care plan discussion, share information with patients consent
- Consider Continuing Health Care funding/DS1500
- Equipment assessment
- Anticipatory medication prescribed and available
- DNACPR considered and discussed, outcome documented, information shared appropriately including ambulance service
- Out of Hours/NWAS updated including DNACPR status and Advance Care Plan
- Referral to other services e.g. Specialist Palliative Care
- Update EPaCCS Record as and when necessary
- ICD discussion and deactivation

### LAST DAYS OF LIFE

**Days**

- Medical review
- All reversible causes of deterioration explored
- Multidisciplinary Team agree patient is in the last days of life
- Clear, sensitive communication with patient and those identified as important to them
- Dying person and agreed others are involved in decisions about treatment and care as they want
- Agree on-going monitoring and support to avert crisis
- Advance Care Planning discussion offered or reviewed
- On-going District Nurse support
- ICD discussion and deactivation if not previously initiated
- Decisions made are regularly reviewed and revised accordingly
- Individual plan of care for the dying person including holistic assessment, review of hydration and nutrition, symptom control etc. is agreed, coordinated and delivered with compassion
- Anticipatory medication prescribed and available to prevent a crisis
- Needs of those identified as important are explored, respected and met as far as possible
- OOH/NWAS updated
- Update EPaCCS Record as and when necessary
- Review package of care if necessary
- Referral to other services e.g. Specialist Palliative Care

### CARE AFTER DEATH

**1 Year/s**

- Nurse verification of death where indicated
- Certification of death
- Clear sensitive communication
- Relatives supported
- Department for Work & Pensions 011 Booklet: What to do after a death or similar
- Post death Significant event analysis
- Update Supportive Care Record/Gold Standards Framework Register/EPaCCS with date and place of death
- Inform all relevant agencies; social care, ambulance service, OOH, Specialist Palliative Care Team, Allied Health Professionals equipment store
- Funeral attendance if applicable and to include carer permission if appropriate
- Follow up bereavement assessment to those identified as important
- Referral of those identified as important to bereavement counselling services as required
- Staff supported

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**ADRT** - Advance Decision to Refuse Treatment  
**DNACPR** - Do Not Attempt Cardio Pulmonary Resuscitation  
**EPaCCS** - Electronic Palliative Care Coordinating System  
**GP** - General Practitioner  
**ICD** - Implantable Cardioverter Defibrillator  
**NWAS** – North West Ambulance Service  
**OOH** – Out of Hours  
**PPC** - Preferred Priorities of Care  

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**Assessment / Description**
Malignant bowel obstruction is a recognised complication of advanced pelvic or abdominal malignancy. May be made worse by adhesions from previous surgery/radiotherapy. Common symptoms associated with malignant bowel obstruction include abdominal pain, abdominal colic, nausea and vomiting. The evidence base for management of malignant bowel obstruction is weak. An individualised approach to management is recommended for each patient and specialist palliative care advice should be sought.

- The diagnosis is made clinically through history and examination
- This may be confirmed with imaging (abdominal X-ray or CT scan) depending on individual circumstances and preferences
- Consider if there are any surgical interventions possible
- Treat constipation if appropriate
- Consider absorption of modified medications when deciding route

**Pharmacology options for Symptom Control in Malignant Bowel Obstruction**

<table>
<thead>
<tr>
<th>Indication(s)</th>
<th>Drug Name</th>
<th>Dose (over 24 hours via CSCI unless otherwise stated)</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Relief of constant pain</td>
<td>Opioid via CSCI/24 hours or transdermal Fentanyl/patch</td>
<td>Dependent on previous dose</td>
<td>Absorption of oral formulation via gut may have been impaired, therefore when converting from oral to CSCI, consider adjusting the dose accordingly.</td>
</tr>
<tr>
<td>Relief of colic</td>
<td>Hyoscine Butylbromide</td>
<td>60mg - 240mg</td>
<td>Do not combine with Cyclizine in CSCI as can cause crystallisation</td>
</tr>
<tr>
<td></td>
<td>Glycopyrronium</td>
<td>600micrograms - 2.4mg</td>
<td>Does not crystallise</td>
</tr>
<tr>
<td>Reduce volume of gastrointestinal secretions</td>
<td>Octreotide</td>
<td>300 - 600micrograms. Doses may be increased up to 1.2mg in some cases under specialist guidance</td>
<td>Can be considered first line. Alternatively use Hyoscine Butylbromide but do not combine with cyclizine in CSCI as can cause crystallisation</td>
</tr>
<tr>
<td></td>
<td>Hyoscine Butylbromide</td>
<td>60mg - 240mg</td>
<td>Do not combine with Cyclizine in CSCI as can cause crystallisation</td>
</tr>
<tr>
<td></td>
<td>Glycopyrronium</td>
<td>600micrograms - 2.4mg</td>
<td>Does not crystallise with other common injectable drugs</td>
</tr>
<tr>
<td></td>
<td>Ranitidine</td>
<td>100mg - 200mg</td>
<td>Does not crystallise with other common injectable drugs</td>
</tr>
<tr>
<td>Reduce tumour oedema. Reduce nausea and vomiting</td>
<td>Dexamethasone</td>
<td>6.6 mg subcutaneously OD or 3.3 mg subcutaneously BD (in morning)</td>
<td>Given as a single dose or divided into 2 doses (before 12 noon). Late administration may cause insomnia/agitation</td>
</tr>
<tr>
<td>Reduce nausea and vomiting</td>
<td>Cyclizine</td>
<td>150mg</td>
<td>Do not combine with Hyoscine Butylbromide in CSCI as can cause crystallisation</td>
</tr>
<tr>
<td></td>
<td>Haloperidol</td>
<td>1.5mg - 5mg</td>
<td>May cause sedation. Use the lowest effective dose. Higher doses may cause sedation.</td>
</tr>
<tr>
<td></td>
<td>Levomepromazine</td>
<td>2.5mg - 25mg</td>
<td>May cause sedation. Use the lowest effective dose. Higher doses may cause sedation.</td>
</tr>
<tr>
<td>Metoclopramide (avoid in complete bowel obstruction)</td>
<td>Metoclopramide</td>
<td>30mg - 60mg. There is an increased risk of neurological adverse effects at doses higher than 30mg/24hour and if used for longer than 5 days.</td>
<td>Contraindicated in complete bowel obstruction. Dose may be increased under specialist Palliative Care advice. Monitor for increased abdominal colic.</td>
</tr>
<tr>
<td></td>
<td>Ondansetron</td>
<td>8mg - 16mg</td>
<td>seek Specialist Palliative Care advice if over 16mg</td>
</tr>
</tbody>
</table>

**Notes**
- Contraindicated in complete bowel obstruction.
- Dose may be increased under specialist Palliative Care advice. Monitor for increased abdominal colic.
- Ondansetron not licenced for SC use.
- Metoclopramide avoids in complete bowel obstruction.
- Dose may be increased under specialist Palliative Care advice.
- Consider absorption of modified medications when deciding route.

**IMPORTANT CONSIDERATIONS:**

**Symptom Control**

**Pain:**
- Opioid analgesia should be titrated to control continuous abdominal pain.
- Colic should be managed with the reduction in dose or discontinuation of prokinetic drugs such as metoclopramide followed by the commencement of an anti-spasmodic such as hyoscine butylbromide

**Reduction of secretions:**
- Patients experiencing large volume vomiting should be prescribed anti-secretory treatment.
- Octreotide is the recommended first line anti-secretory medication

**Reduction of nausea and vomiting:**
- Anti-emetics should be administered via the subcutaneous route. Prokinetics are not advised in a bowel obstruction affecting the small bowel or in a complete obstruction at lower levels of the bowel

**Corticosteroids:**
- A five day trial of Dexamethasone 8mg daily orally or similar dose, subcutaneously should be considered in all patients to reduce tumour related oedema

**Laxatives:**
- The use of stimulant laxatives should be avoided. The use of stool softeners may be appropriate

**Interventions**

**Medication Delivery:**
- Medication should be delivered via the subcutaneous route due to potential problems with absorption

**Nasogastric Tubes:**
- A wide bore nasogastric tube should be considered for patients with upper gastrointestinal obstruction or large volume vomiting

**Venting Gastrostomies:**
- Venting gastrostomies or jejuno stomies should be considered for patients with malignant bowel obstruction who have a prognosis of greater than 2 weeks.
- Venting gastronomies have been shown to be cost effective with low morbidity and mortality
**Assessment / Description**

Causes of breathlessness can be multi-factorial: physical, psychological, social and spiritual factors can all contribute to a person feeling breathless. Assessment is vital, particularly in a new presentation. Undertake a history and clinical examination, including oxygen saturations. Investigations such as chest x-ray may be necessary and management will depend on clinical diagnosis. Treat what may be caused by an acute event and reversed, e.g. infection, anaemia, pulmonary oedema etc.

**Pharmacological Options**

**Opioids**: Morphine oral solution 2.5mg QDS regularly and 4 hourly PRN. The maximum dose of oral Morphine that is likely to be helpful for breathlessness is 30mg/24 hour period.

If patient is unable to tolerate oral meds, subcutaneous Morphine Sulphate – 2.5mg 4 hourly prn (1.25mg in opioid naïve patients)

If the patient’s eGFR is <30mls/min an alternative opioid should be considered. There is currently no evidence for the use of alternative opioids in the management of breathlessness.

**Oxygen**: The evidence for efficacy is limited.

A trial of oxygen should be considered in patients known to have O2 saturation less than 90%. PRN use of Oxygen should be avoided.

Considerable care should be taken in patients with known COPD/Type 2 respiratory failure - watching for CO2 retention headache, flushed skin, fast pulse, hand flap, drowsiness etc.

**Corticosteroids**: may help in patients with tumour compression or lymphangitis carcinomatosis.

No evidence of benefit in non specific dyspnoea.

Dose: (before 2pm)

Dexamethasone 4mg – 8mg daily PO
Or 8mg – 16mg PO in lymphangitis or Superior Vena Caval Obstruction (SVCO)

**Nebulised Medication**: Sodium Chloride 0.9% may help as a mucolytic, 2.5 - 5ml 4hrly PRN

Consider a bronchodilator for bronchospasm e.g. Salbutamol 2.5mg 6hrly PRN (may be used more frequently in some cases)

**Benzodiazepines**: Lorazepam 500micrograms - 1mg SL PRN 2- 4 hourly (max dose 4mg in 24 hour period).

If patient unable to tolerate oral medication, consider subcutaneous Midazolam 2.5mg - 5mg 4hrly prn.

If effective this can be incorporated into a syringe pump over 24 hours

Be careful of the risk of both dependence and tolerance if used for the more than 7 days.

**Non-Pharmacological options for managing breathlessness**

- Calm Environment
- Acknowledgment and explanation
- Adequate positioning of the patient to aid breathing
- Use of fan or cool air across face
- Breathing exercises and relaxation training
- Acupuncture, aromatherapy and reflexology

**As condition improves, reduce monitoring and evaluate treatment and stop interventions that are no longer needed**

**If breathlessness persists and causes distress consider appropriate pharmacological options**

**Treat reversible causes of breathlessness and & monitor response**

PLUS
Start appropriate non pharmacological interventions (blue boxes)

**If breathlessness persists and causes distress consider appropriate pharmacological options**

(purple boxes)

As condition improves, reduce monitoring and evaluate treatment and stop interventions that are no longer needed
Assessment/Description

Constipation is defined by the patient and is a symptom not a disease. The cause of the constipation should be identified and treated, managing bowel obstruction where appropriate. Aim to prevent constipation by the early introduction of laxatives, especially if patients are taking pain killers regularly.

- History, normal bowel habit, medicines other causative factors
- Abdominal palpation and auscultation and digital rectal examination
- Investigation abdominal x-ray, check calcium levels
- Treatment should be individualised to the patient and what they are able to tolerate. In most cases the oral route to manage constipation should be used initially. If constipation is not resolved after 5-7 days seek Specialist advice

Quick guide: Constipation

Reference: Merseyside and Cheshire Palliative Care Network Audit Group Guidelines on Constipation

### Causes to consider:

- Drug-induced including opioids, diuretics, anti-cholinergics, ondansetron, chemotherapy
- Dehydration
  - Review diuretics and fluid intakes
- Reduced mobility
- Hypercalcaemia - consider IV fluids and bisphosphonates
- Environmental - lack of privacy
- Concurrent disease
- Altered dietary intake - increase fluid and fibre intake if possible
- Neurological
- Intestinal obstruction

### Treatment and management

<table>
<thead>
<tr>
<th>Oral laxatives commonly used in palliative care</th>
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<tbody>
<tr>
<td><strong>Type of laxative</strong></td>
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<tr>
<td>Stimulant laxatives</td>
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<tr>
<td></td>
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<td></td>
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<tr>
<td></td>
</tr>
<tr>
<td>Combination laxatives</td>
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<tr>
<td></td>
</tr>
<tr>
<td>Osmotic laxatives</td>
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<td></td>
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<tr>
<td>Opioid induced constipation</td>
</tr>
</tbody>
</table>

### Rectal interventions for constipation

- For patients with established constipation, it is usually most effective to combine faecal softeners and stimulant laxative. If necessary, an osmotic agent can then be added on a pm or regular basis
- Oral laxatives should be reviewed every 3 to 4 days using stool consistency chart (e.g. Bristol stool chart)
- The use of rectal interventions should be guided by the findings on rectal examination
- Enemas including phosphate and sodium citrate versions - follow local guidance

<table>
<thead>
<tr>
<th>Rectal Intervention</th>
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</thead>
<tbody>
<tr>
<td>Impacted hard faeces</td>
</tr>
<tr>
<td>Bisacodyl 10mg Suppository, plus glycerol 4g suppository (plus combination oral laxative)</td>
</tr>
<tr>
<td>If ineffective use enema</td>
</tr>
<tr>
<td>Empty rectum plus loaded colon</td>
</tr>
<tr>
<td>Phosphate enema (plus combination oral laxative)</td>
</tr>
<tr>
<td>Impacted soft faeces</td>
</tr>
<tr>
<td>Bisacodyl 10mg Suppository (plus oral stimulant)</td>
</tr>
<tr>
<td>If ineffective use enema</td>
</tr>
</tbody>
</table>
QUICK GUIDE NAUSEA & VOMITING (FOR LAST DAYS OF LIFE - SEE PAGE 17)

Reference Merseyside and Cheshire Palliative Care Network Audit Group Guidelines on Nausea and Vomiting

Assess the likely cause for nausea to guide the anti-emetic most likely to relieve symptoms

Review reversible causes (see boxes below)

Initial Treatment

Patients who become nauseated or start vomiting:

Gut causes

Metoclopramide 10mg TDS PO/SC or CSCI 30mg/24h (avoid in complete bowel obstruction—see guidance on bowel obstruction). There is an increased risk of neurological adverse effects at doses higher than 30mg/24h and if used for longer than 5 days.

Domperidone 10mg TDS PO. There is an increased risk of cardiac side effects at dose higher than 30mg/24h and if used for longer than 7 days. — see BNF for more information

Non gut causes (e.g. medication, renal failure, biochemical disturbances, or cerebral causes)

Metoclopramide 10mg TDS PO/SC or 30mg CSCI—see warning above

Haloperidol 500 micrograms–3mg PO/SC at night or CSCI 1.5mg to 5mg

Cyclizine 50mg TDS PO/SC or CSCI 150mg over 24 hours in water for injection

Levomepromazine 6mg PO or 2.5mg SC at night or CSCI 6.25mg-12.5mg (can use 3mg PO if sedation a problem)

Alternative anti-emetics may be more appropriate in certain circumstances

1. Severe Heart Failure:
Levomepromazine 6mg PO or 2.5mg SC at night or CSCI 6.25mg-12.5mg (can use 3mg PO if sedation a problem)

Avoid anti-emetics with anti-muscarinic side effects, such as Cyclizine, that may cause tachy-arrthymias.

2. Toxicity or metabolic or biochemical cause of vomiting (medication related, renal failure, uraemia, hypercalcaemia)

Haloperidol 500 micrograms - 3mg PO/SC at night or CSCI 1.5mg to 5mg

Cyclizine 50mg TDS PO/SC or CSCI 150mg over 24 hours in water for injection

Levomepromazine 6mg PO or 2.5mg SC at night or CSCI 6.25-12.5mg (can use 3mg PO if sedation a problem)

3. Parkinson’s Disease / Lewy Body Dementia:

Avoid anti-emetics with a dopamine receptor antagonist effect e.g. Haloperidol, Levomepromazine and Metoclopramide.

Domperidone 10mg TDS PO first line - see caution above

Ondansetron 4mg PO/SC PRN can be considered.

4. Raised Intracranial Pressure (ICP):

If taking oral Dexamethasone for symptoms of raised ICP, this should continue to be given daily via the SC route.

Aim to maintain at the lowest maintenance dose that controls the symptoms of raised intracranial pressure.

Dexamethasone subcutaneously 3.3mg to 16.5mg max daily dose in 1 or 2 divided doses.

All doses of dexamethasone should be given before 2pm

*Dexamethasone can raise blood sugar levels and capillary blood glucose levels should be checked as per local guidance

If there is a risk of seizures, e.g. in brain metastasis, avoid the use of levomepromazine which can lower the seizure threshold

5. Gastric Outlet Obstruction / Reflux:

This can occur due to autonomic neuropathy (paraneoplastic) medication (anti-cholinergics), metabolic causes (e.g. hypercalcaemia), mechanical obstruction.

If complete bowel obstruction is excluded:

Metoclopramide 30mg in CSCI over 24 hours - higher doses can be used - seek specialist advice

Avoid the concurrent prescribing of a prokinetic e.g. metoclopramide and an anti-cholinergic (e.g. Cyclizine) which will inhibit it’s prokinetic action. Metoclopramide can cause colic.

6. Bowel Obstruction:

See guidance on bowel obstruction - page 7
### COMMON TYPES OF PAIN

<table>
<thead>
<tr>
<th>Visceral / Soft Tissue Pain (nociceptive)</th>
<th>Bone Pain (somatic nociceptive)</th>
<th>Nerve Pain (neuropathic)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Constant dull pain; Poorly localised</td>
<td>Usually well localised; worse on movement; localised tenderness</td>
<td>Try opioids first, but may be less responsive. Consider adjuvant neuropathic analgesia</td>
</tr>
</tbody>
</table>

#### WHO STEP 1: Non-Opioids
- e.g. Paracetamol 1g qds PO
  +/− ADJUVANT

#### WHO STEP 2: Non-Opioid plus Weak Opioid
- e.g. Codeine 30-60mg qds PO
  +/− ADJUVANT

#### WHO STEP 3: Non-Opioid plus Strong Opioid
- e.g. Morphine
  +/− ADJUVANT

#### USE OF TRANSDERMAL OPIOID PATCHES
- Commencing transdermal opioid (patches):
  - Titrate with 4 hourly immediate release oral Morphine, until pain is controlled
  - Remember a Fentanyl 25micrograms/hour patch is equivalent to a 60-90mg total daily dose of oral Morphine
  - Stick patch to hairless skin; clip (do not shave) hair
  - Initial analgesic effect will take at least 12-24 hours, and a steady state may not be achieved for 72 hours
  - Ensure immediate release oral Morphine (or alternative) is available for breakthrough pain
  - Change patch every 72 hours; use a new area of skin
  - A 12-24 hour depot of drug remains in the patch when removed; fold in on themselves and discard safely out of the reach of children / vulnerable adults
  - Opioid withdrawal may occur when switching from Morphine to Fentanyl; manage with PRN Morphine.

### Clinical Practice Summary

**Pain can be improved for patients. If not improving, seek Specialist Palliative Care advice**

“Dose adjustments may need to be made for renal failure” - See renal failure page 20

### Conventional Opioid Titration

**IMMEDIATE RELEASE MORPHINE**
- (4 hourly duration of action)
- Regularly: Morphine Oral Solution 2.5mg - 5mg 4 hourly
- PRN: Morphine Oral Solution 2.5mg - 5mg 2 hourly
- If clinically frail or eGFR less than 60ml/min use lower doses or reduced frequency of dose e.g. 6 or 8 hourly.
- Assess response of background pain to opioids and if necessary increase dose by 30-50% every 24-48 hours to achieve pain control. If not – seek Specialist Palliative Care advice.
- If eGFR less than 30ml/min see renal failure page 20

**SUSTAINED RELEASE MORPHINE**
- (12 hourly preparation)
- Zomorph capsules BD, MST tablets BD, Morphgesic SR BD, Filnarine SR BD
- e.g. 5mg morphine used 4 times = 20mg oral morphine in 24 hours = 10mg sustained release morphine (12 hourly) twice a day

### Alternative Opioid Titration

**SUSTAINED RELEASE MORPHINE**
- (12 hourly duration of action)
- Regularly: Morphine MR 10mg BD 12 hourly
- Zomorph capsules , MST tablets , Morphgesic MR Filnarine SR
- PRN: Morphine oral solution 2.5 - 5mg 2 hourly
- Assess response of background pain to opioids and if necessary, increase dose by 30 - 50% every 24-48 hours to achieve effective breakthrough dose – consider Co-analgesics.
- If clinically frail or eGFR less than 60ml/min use modified release medication with caution.
- If eGFR less than 30ml/min see renal failure page 20
- When pain controlled calculate total daily dose of modified release morphine and any immediate release morphine taken in a 24hour period and divide by 2 to get a 12 hourly dose.
- Always ensure starting rescue breakthrough dose is around 1/6th of the total 24 hour dose

### CO-ANALGESICS
- • Neuropathic Pain Agents
  - Gabapentin start 100mg to 300mg nocte
  - Pregabalin start 25mg OD or BD
  - Amitriptyline 10mg nocte
  (starting doses in clinical frailty, requires titration to effect)
- • NSAIDS (Ibuprofen 400mg TDS or Naproxen 500mg BD) with food
  - See page 12 for more detail

### Anticipate opioid side effects
- Always co-prescribe regular laxatives
  - Senna or Docusate or Co-danthramer or Macrogol
  - and consider PRN
  - Anti-emetics such as
  - Metoclopramide 10mg TDS PO
  - Or Haloperidol 500 micrograms to 3mg PO at teatime
  - Or Cyclizine 50mg TDS PO or Levomepromazine 3 to 6mg PO nocte

### Guidance in the Last Days of Life (see page 17)
- • When a patient is in the dying phase, LEAVE PATCH IN SITU, and change regularly as before.
  - If patient has pain use an appropriate subcutaneous dose of opioid PRN for breakthrough pain
  - If PRN doses are needed more that twice start CSCI in addition to patch
  - Ensure PRN dose adequate for both patch & CSCI
  - Seek Specialist Palliative Care advice for support if needed.
Clinical Practice Summary
Lancashire and South Cumbria Consensus Guidance - August 2017

Quick Guide - Pain (Part 2) - Complex Pain

Reference
Merseyside and Cheshire Palliative Care Network Audit Group Guidelines on Neuropathic Pain, Symptom Control Medication in the Dying Person, Opioid Substitution, Transdermal Opioids in Palliative Care

If eGFR less than 30ml/min see RENAL FAILURE Page 20

Pain can be improved for patients. If not improving; seek Specialist Palliative Care advice, especially if:
- Complex, multiple pains where assessment is difficult
- Pain appears to be resistant to usual measures or not responding to Morphine doses equivalent to or exceeding 120mg Morphine in 24 hours
- Difficulty in managing pain due to adverse effects of medication or compliance

Concept of Total Pain
Should prompt healthcare professionals to consider all possible influences on the individual’s pain experience:
- Physical
- Spiritual
- Social
- Psychological

Success in pain management depends on:
- Regular review of the pain and its causes
- Effectiveness of treatment
- Acceptability of the proposed treatment to the patient

The patient’s understanding, fears, concerns and previous experience of pain, as well as their expectations of treatment will all influence each individual’s experience of pain and its effective management.

A Guide to Equivalent Doses of Opioid Drugs
This table of doses is a guide - not a set of definitive equivalences.

Use the table to identify an appropriate starting point for your prescribing decision. Ask if the pain is opioid responsive. All prescribing decisions must be based on a full clinical assessment. Higher opioid doses may be needed for some patients - seek advice

Think about the role of adjuvant medication before rotating opioids, changing the dose or route.

Consider reducing prescribed opioid dose by 30-50% if converting from one route to another route (e.g. transdermal to oral or oral to subcutaneous) or there is concern about opioid toxicity (confusion, drowsiness, myoclonic jerks, slowed respiration, pin-point pupils)

Be aware of drug interactions and remember individual patients may metabolise different drugs at varying rates.

Never increase an opioid dose by more than 50% of the previous 24 hour regular dose without SPECIALIST ADVICE

<table>
<thead>
<tr>
<th>Oral Morphine</th>
<th>Oral Oxycodone</th>
<th>Transdermal Buprenorphine</th>
<th>Transdermal Fentanyl</th>
<th>Subcutaneous Morphine</th>
<th>Subcutaneous Oxycodone</th>
</tr>
</thead>
<tbody>
<tr>
<td>4 hr dose (mg)</td>
<td>12 hourly dose (mg)</td>
<td>4 hr dose (mg)</td>
<td>12 hourly dose (mg)</td>
<td>4 hr dose (mg)</td>
<td>24 hr CSCI dose (mg)</td>
</tr>
<tr>
<td>(break-through dose)</td>
<td></td>
<td>(break-through dose)</td>
<td></td>
<td></td>
<td>(break-through dose)</td>
</tr>
<tr>
<td>1.25</td>
<td>5</td>
<td>-</td>
<td>-</td>
<td>5</td>
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<td>2.5</td>
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<tr>
<td>20</td>
<td>60</td>
<td>15</td>
<td>45</td>
<td>-</td>
<td>52.5</td>
</tr>
</tbody>
</table>

**SEEK SPECIALIST ADVICE**

| 30 | 90 | 20 | 60 |
| 40 | 120 | 25 | 75 |
| 50 | 150 | 30 | 90 |
| 60 | 180 | 40 | 120 |

**SEEK SPECIALIST ADVICE**

| 70 | 75 |
| 105 | 100 |
| 122.5 | 125 |
| 140 | 150 |
| 15 | 90 | 10 | 60 |
| 20 | 120 | 12.5 | 75 |
| 25 | 150 | 15 | 90 |
| 30 | 180 | 20 | 120 |

Neuropathic Pain Agents

AMITRIPTYLINE - start 10mg OD increased to 25mg OD after 3-7 days and then by 25mg every 1 - 2 weeks to a maximum of 75mg daily

GABAPENTIN - start 100mg OD increase to 100mg BD after 2-3 days to 100mg TDS after 2-3 days and then by increments of 100mg every 2-3 days depending on response to a maximum dose of 900mg TDS

PREGABALIN - start 25mg BD and increase by 25mg every 2-3 days to a maximum dose of 300mg BD

DULOXETINE - start at 30mg OD and increase to 60mg OD after 2 weeks - stop if no response after 2 months. Maximum dose 120mg OD

Start with either an anticonvulsant or an antidepressant and titrate dose as above. Response takes a number of days to become apparent. If no apparent response seek advice from Specialist Palliative Care team.

NB Prescribing of the above for some types of neuropathic pain is beyond licence. The prescriber should follow relevant professional guidance, taking full responsibility for the decision.

Informed consent should be obtained and documented.
HYPERCALCAEMIA

• Hypercalcaemia is common in cancer of breast, myeloma, lung, head and neck, kidney, thyroid and cervix.

• Primary hyperparathyroidism should be considered as a possible cause (6% of cancer patients)

Presentation:
• Symptoms of hypercalcaemia include: fatigue, weakness, constipation, nausea, vomiting, polyuria, polydipsia, cardiac arrhythmias, delirium, drowsiness and coma.

• Corrected serum calcium >2.7mmol/L (some variation between laboratories)

ASSESSMENT:
Clinical assessment of the patient is crucial in determining whether treatment of hypercalcaemia is appropriate, as it generally requires IV fluids and admission to an institution.

Generally a decision to treat should be motivated by the patient’s symptomatology rather than absolute calcium level. The most important goal of treatment is to improve clinical symptoms. Hypercalcaemia may be a poor prognostic sign in cancers such as lung and cervix.

Onset of symptoms raising clinical suspicion should be investigated. Bloods should be checked for urea and electrolytes (U&Es), estimated glomerular filtration rate (eGFR), liver function tests (LFT’s) and calcium.

TREATMENT:
May require in-patient unit care in hospital or hospice. (Refer to local guidelines around bisphosphonate dosing)

The patient should be rehydrated with 1-3 litres of parenteral 0.9% sodium chloride before the administration of bisphosphonates. The volume and rate of fluid replacement should be adjusted in each patient according to their age, rather than absolute calcium level. The most important goal of treatment is to improve clinical symptoms. Hypercalcaemia may be a poor prognostic sign in cancers such as lung and cervix.

The treatment of choice after rehydration is intravenous bisphosphonate - pamidronate, zoledronic acid or ibandronate depending on local formulary choices.

Corrected calcium levels should be rechecked at 5-7 days after the bisphosphonate infusion. Checking calcium levels prior to this is not appropriate, as the bisphosphonate will not have achieved it’s maximal effect.

• Consider Advance Care Plan about how and where to manage further episodes in the future.

EPILEPTIC SEIZURES

• May settle spontaneously

• Ensure airway secure and administer oxygen if available

• If seizure does not stop within 5 minutes give either
  - Subcutaneous, intranasal or buccal midazolam 5mg to 10mg OR
  - Diazepam 10mg-20mg rectally

Once settled consider long term seizure management with relevant specialists if not in last weeks/days of life. Alternative medication may be considered - please see local guidance.

IF SEIZURES CONTINUE despite above measures for a further 5 minutes - Repeat measures above

• Decide if transfer to hospital for emergency management is needed or if care will continue in the current care setting

• For acute management - a secure airway should be established, oxygen should be administered, cardiorespiratory function should be assessed and intravenous access should be established.

• If patient is to stay at home or hospice and two doses needed, consider a continuous subcutaneous infusion of Midazolam 20-30mg over 24 hours, seek Specialist advice.

SUPERIOR VENA CAVAL OBSTRUCTION (SVCO)

• Compression / invasion or thrombosis of SVC due to tumour or nodal mass within mediastinum, preventing venous drainage from head, arms and upper trunk

• Commonest causes (95%) – lung cancer, non-Hodgkin’s lymphoma

• Usually onset over weeks or months, but occasionally occurs rapidly over days

MANAGEMENT:
Administer Dexamethasone 16mg orally or parenterally in one or two divided doses -IMMEDIATELY URGENTLY (ideally the same day) discuss with Oncologist about future management

Consider the possibility of radiological stenting.

SYMPTOMS/SIGNS:
• Swelling of face, neck, arms
• Headache
• Dizziness/ Visual disturbance
• CNS depression
• Seizures

• Dyspnoea
• Dilated veins – neck, trunk, arms
• Hoarse voice
• Stridor
• Cyanosis
### Quick Guide: Palliative Care Emergencies - Part 2

**Reference**
Merseyside and Cheshire Palliative Care Network Audit Group Guidelines on Metastatic Spinal Cord Compression Major Haemorrhage

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### Metastatic Spinal Cord Compression

- Affects 5-10% of patients with cancer
- Most common in prostate, lung, breast cancer and myeloma
- Catastrophic event – aim is to prevent establishment of permanent loss of function
- Symptoms may be vague, there should be a high index of suspicion if a patient goes “off their legs”, becomes unsteady, struggles to get out of a chair or climb stairs.
- Patients with cancer and neurological signs or symptoms of spinal cord compression should be treated as an oncological emergency

**Follow Local Oncology Guidance**

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### Same Day - Medical Assessment

Full history and neurological examination, Assess fitness to treat

**Same Day - Contact**: -
METASTATIC SPINAL CORD CO-ORDINATOR at Oncology centre to discuss case (for Lancashire and South Cumbria 01772 71656 Or Bleep 2664)

### If Suspected:

- Give Dexamethasone 16mg BY MOUTH or convert to SC
- Prescribe medication for gastric protection
- Give adequate analgesia (opioid if necessary) to enable transfer for admission / investigation
- Nurse flat if pain / symptoms suggest spinal instability
- Request urgent admission and MRI scan

### Post Diagnosis

May have radiotherapy or spinal surgery to stabilise spine and relieve pressure on spinal cord. Aim to maintain function and continence as much as possible. Involve physiotherapy and occupational therapy as soon as possible. Titrate steroids down to the lowest dose over 2 - 4 weeks dependent on patient’s symptoms and condition. In many cases developing metastatic spinal cord compression is a poor prognostic sign

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### Major Haemorrhage

**Clinical Presentation**:
- Cardiovascular compromise – hypotension, tachycardia (>100bpm = significant recent bleed)
- Identifiable bleeding source – haematemesis, haemoptysis, PV or PR bleeding, haematuria, melena
- Erosion of an artery by a malignant ulcer or superficial/fungating tumour
- Bleeding of all types occurs in 14% of patients with advanced disease - seek Specialist advice if time and clinical situation permit
- Haemorrhage causes death in approximately 6% patients
- Catastrophic external haemorrhage less common than internal bleeding
- It may be a terminal event in both advanced cancer and non-malignant disease.

**Management**:

**A member of staff must remain with the patient to provide support at all times**
- Plan ahead where possible, record and share information with key organisations via EPaCCS
- If there are warning signs or high anticipated risk of bleeding have a proposed management plan ideally discussed with patient and/or family and staff
- Record management plan in case notes and communicate this to all team members
- Provide dark coloured towel to disguise blood loss.
- Anticipatory prescribing of Midazolam 10mg IM, SC, buccal or sublingual.
- The subcutaneous route may be less effective in catastrophic bleeds due to peripheral shut down with unpredictable absorption of the medication

**Further Care**:
It may be necessary to commence and continue an infusion of anxiolytic (Midazolam) and/or analgesic e.g. Morphine or Oxycodeine) in the last hours of life. If bleeding temporarily stops further management will depend on overall clinical status and discussion with patient and family in relation to further acute interventions. Consider referral for diathermy, radiotherapy or embolisation depending on local availability and protocols. Seek specialist palliative care advice around other options to prevent re-bleeding

---

### Symptoms – particularly new or changing:

**Back/Spinal Pain**:
- may radiate in a radicular, ‘band-like’ pattern
- progressive / unremitting
- may be worse on coughing or straining
- may be nocturnal, pain preventing sleep
- may not be present

Nerve root pain in limbs.
Weakness of limbs (out of proportion to general condition of patient)
Difficulty walking.
Sensory changes - tingling, numbness, “my legs don’t belong to me.” Difficulty passing urine – usually a late presentation.
Constipation or faecal incontinence.

**Signs: Do not wait for signs. Act on the symptoms**

Localised spinal tenderness
Weakness of limbs
Reflexes: Absent / increased. Extensor plantars.
Altered sensation - look for a sensory level
Distended bladder

Contact local Specialist Palliative Care Team if advice on symptom management required
QUICK GUIDE  |  CARE IN THE LAST WEEKS OR DAYS OF LIFE
Reference: Merseyside and Cheshire Palliative Care Network Audit Group Guidelines on Symptom Control in the Dying Person

FIVE KEY PRIORITIES

RECOGNISE:
• The possibility that a person is in the last weeks of life or they may die within the next few days or hours and communicate this clearly:
• Consider and address reversible causes where appropriate / possible
• Identify and where possible make decisions in accordance with the individual’s wishes and needs
• Review the assessment and decisions on a regular basis

COMMUNICATE:
• SENSITIVELY with the individual and those important to them

INVOLVE:
• All relevant people in making decisions as far as they indicate they want to be

SUPPORT:
• The family and other people important to the dying person by exploring, respecting and meeting their needs where possible

PLAN:
• Create an individualised plan of care. This should include decisions around:
  - Cardiopulmonary resuscitation
  - Facilitating or preventing change in place of care
  - Supporting oral food and fluid intake
  - Stopping or continuing physical observations and / or investigations
  - Starting, stopping or continuing clinically assisted hydration and / or nutrition
  - Review of long term medication - stopping those that are no longer needed and switching others to a route which ensures they can continue and provide benefit
  - Anticipatory prescribing of medication for the common symptoms at end of life (i.e. pain, breathlessness, respiratory tract secretions, agitation, nausea and vomiting) and other problems specific to that individual such as management of seizures or bleeding, etc.

QUICK GUIDE  |  DIABETES MANAGEMENT IN THE LAST WEEKS OF LIFE
Reference: Northern England Strategic Clinical Network Guidance

Assessment/Description
Explore with the individual and those important to them changing the approach to diabetes management including:
• The aim of management - avoiding hypoglycaemia rather than avoiding longer term complications due to hyperglycaemia
• The value of continuing to monitor blood glucose readings
• The method and frequency of checking blood glucose levels
• The type of management - tablets and / or insulin

Devise a management plan with the patient and those important to them. Ensure your local diabetes specialist team are involved if the patient remains on insulin. Aim to:
• Keep invasive tests to a minimum
• Be alert to symptoms that may be due to hypo or hyperglycaemia and have appropriate medication / interventions available to address these if they develop

Aim for a Target BM reading between 6 and 15.

Type 2 Diabetes
Diet controlled

Stop monitoring blood sugars on a regular basis

If insulin stopped:
 Urinalysis for glucose
 If over 2+ positive check capillary blood glucose.
 If over 20mmol/l give 6 units of Actrapid insulin
 Recheck blood sugar level after 2 hours

If blood sugar still high start long-acting insulin
 Glargine Lantus

Type 2 Diabetes
On tablets and / or insulin

Stop oral hypoglycaemics
 Consider stopping insulin depending on dose

Either:
 If insulin is to continue:
 Prescribe once daily long-acting insulin e.g. insulin Glargine Lantus in the MORNING with a 25% reduction in the daily insulin dose

Type 1 Diabetes
On regular insulin

Continue once daily long-acting insulin in MORNING e.g. insulin Glargine Lantus with a 25% reduction in dose
Stop short-acting insulin

Check blood sugars once daily at TEA TIME
 If below 6mmol/l reduce insulin dose by 25%
 If above 15mmol/l increase dose of insulin to reduce risk of ketosis
 After dose by 2 units if daily dose of insulin is below 50 units or more
 After dose by 4 units if daily dose is 50 units or more
## Quick Guide

<table>
<thead>
<tr>
<th>CONTINUOUS SUBCUTANEOUS INFUSIONS (CSCI) / Syringe Pump - also see local guidelines</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reference</td>
</tr>
</tbody>
</table>

### Assessment/Description

Syringe PUMPS are used to administer medication by a continuous subcutaneous infusion (CSCI) over a 24 hours period. They are classed as high risk devices and should only be used by suitably trained clinicians.

### Indications for commencing medication via continuous subcutaneous infusion (CSCI)

- Patient is unable to take oral medication due to:
  - Nausea and vomiting
  - Difficulty in swallowing
  - Intestinal obstruction
- Malabsorption / uncertain absorption of oral medication
- For care in last days of life when oral route is unreliable and regular medication is needed to maintain comfort

### Diluent

Most commonly used medication in a syringe pump should be diluted with water for injection. Drugs may be diluted with Saline 0.9% except Cyclizine or Diamorphine (doses above 40mg) which should be diluted in Water for injection.

### Syringe Pump

All syringe pumps must be serviced regularly according to local guidance and at least annually, whether used or not to ensure their function is maintained. Syringe pumps should be sent for maintenance checks immediately if they have been dropped, suffered fluid ingress (e.g. had fluid spilt over them or dropped in a bath) or if there is any doubt as to their functional operation whilst in use.

The following points should be taken into account when using syringe pumps:

- Protect the syringe from direct sunlight whenever possible
- Carry out a visual inspection of the solution within the syringe at each monitoring (refer to local policy) check and discard if evidence of crystallisation or precipitation, cloudiness or change in consistency
- Avoid mixing medicines in one syringe if compatibility data is not available; do not mix more than three medicines unless on the advice of a palliative care Specialist

### Syringe Pump site selection:

- Oedematous areas including lymphoedematous arms (poor drug absorption, and increased risk of infection/exacerbation of oedema)
- Bony prominences (poor absorption and discomfort)
- Irradiated sites (may have poor perfusion and hence poor drug absorption)
- Skin folds, sites near a joint and waistband area (movement may displace cannula or cause discomfort)
- Broken skin
**QUICK GUIDE**

**PAIN IN THE LAST DAYS OF LIFE**

Reference: Merseyside and Cheshire Palliative Care Network Audit Group Guidelines on Symptom Control in the Dying Person

**GENERAL COMMENTS**

In the majority of cases injectable Morphine is the first line opioid of choice in the last days of life.

If patient has been well established on an alternative opioid such as Oxycodone continue it and follow the principles outlined in the flow diagrams.

For patients who have not previously been given medicines for pain management, start with the lowest effective dose of pain killer and titrate as clinically indicated.

Alternative opioids may be needed if the patient has significant renal impairment - seek specialist advice.

**Non-Pharmacological options**
- Reposition patient
- Heat / cold packs
- Distraction
- Acknowledgement and explanation

**Patient is in pain**

Is patient already taking oral morphine?

- **YES**
  - Continue oral morphine if patient still has a safe swallow
  - When swallow lost: Convert to a continuous subcutaneous infusion (CSCI) by adding up the total oral dose over 24 hour and dividing by 2 (taking into account additional rescue doses that have been taken in the last 24hours)
  - Ask for advice from Specialist Palliative Care team or Pharmacy, if unsure or calculated dose is above 60mg /24hour via CSCI
  - Prescribe: PRN rescue dose of Morphine around 1/6th of the 24hour dose in the CSCI 2hourly PRN
  - After 24hours review.
  - If three or more doses required PRN consider a continuous subcutaneous infusion over 24 hours

- **NO**
  - Continue oral morphine if patient still has a safe swallow

**Patient’s pain is controlled**

Is patient already taking oral morphine?

- **YES**
  - When swallow lost:
    - Convert to a continuous subcutaneous infusion (CSCI) by adding up the total oral dose over 24 hour and dividing by 2 (taking into account additional rescue doses that have been taken in the last 24hours)
    - Ask for advice from Specialist Palliative Care team or Pharmacy if unsure or calculated dose is above 60mg/24hour via CSCI
  - Prescribe:
    - PO Morphine 5mg or SC Morphine 2.5mg 1 hourly PRN
    - PO Morphine 2.5mg or SC Morphine 1.25mg 2 hourly PRN if frail or renal impairment
    - If requires three or more doses within 4 hours seek medical review and increase PRN dose to 10mg PO or 5mg SC 2hourly
  - After 24hours review
  - If three or more doses required PRN consider a continuous subcutaneous infusion over 24 hours

- **NO**
  - Prescribe: PO Morphine 5mg or SC Morphine 2.5mg 1 hourly PRN
  - PO Morphine 2.5mg or SC Morphine 1.25mg 2 hourly PRN if frail or renal impairment
  - If requires three or more doses within 4 hours seek medical review and increase PRN dose to 10mg PO or 5mg SC 2hourly

**ADDITIONAL INFORMATION**

**Transdermal opioid patches at end of life (Fentanyl /Buprenorphine)**

It is recommended that opioid patches are left in place and changed regularly in last days of life.

If pain occurs a rescue dose of an appropriate injectable opioid is administered - see page 10 for guidance about equivalent doses.

If 2 or more rescue doses are needed in 24hours consider setting up a CSCI with the total dose of rescue medication given in the previous 24 hours up to a maximum of 50% of the existing regular opioid (patch) dose.

Remember to combine the dose of the opioid patch and the dose of opioid in the syringe pump to work out the new rescue dose (1/6th - 1/10th of the total 24hour dose)

IF YOU ARE IN ANY DOUBT ABOUT HOW TO MANAGE A PATIENT’S PAIN IN THE LAST DAYS OF LIFE ASK FOR SPECIALIST ADVICE
# QUICK GUIDE RESPIRATORY TRACT SECRETIONS IN THE LAST DAYS OF LIFE

## Reference
Merseyside and Cheshire Palliative Care Network Audit Group Guidelines on Symptom Control in the Dying Person

## Assessment/Description
Patient unable to clear secretions from their upper respiratory tract properly, causing secretions to move as they breathe, creating noise

## Non-Pharmacological options
- Reposition patient
- Active surveillance
- Acknowledgement and explanation
- Cautious suction

## Pharmacological Options:

**INITIALLY:**
- Glycopyronium 200 micrograms SC PRN 2 hourly
- Or (alternative) Hyoscine Butylbromide 400micrograms SC PRN 2 hourly

**ONGOING:**
- Anti-cholinergics
  - **Dose Range via CSCI/24 hours**
    - Glycopyronium: 600 micrograms to 1.2mg
    - Hyoscine Butylbromide: 60mg to 240mg
    - Hyoscine Hydrobromide: 1.2mg to 2.4mg
  - In significant renal impairment use Glycopyronium
  - Use Hyoscine Butylbromide with caution with patients with cardiac failure (risk may not be relevant in last days of life)
  - Seek Specialist Palliative Care advice if patient not settling

## Assessment/Description

**Non-Pharmacological options**

- Reposition patient
- Active surveillance
- Acknowledgement and explanation
- Cautious suction

**Pharmacological Options:**

**INITIALLY:**
- Midazolam 2.5mgs - 5 mgs SC up to 2 hourly PRN.
  - If eGFR < 30 give a reduced dose of Midazolam e.g. 1 mg to 2.5 mgs SC PRN

**ONGOING:**
- **Anti-cholinergic side effects can arise:** treat this with frequent mouth care which may include artificial saliva replacement gels or sprays.
- Secretions which have already accumulated will not be removed by medication. Early treatment improves the prospect of achieving symptom control.
- If one agent doesn’t work, try switching to the other after full titration to maximum dose over 24 hours; if there is still no improvement, consider stopping medication.
- Seek Specialist advice as required.
- Hyoscine Hydrobromide crosses the blood brain barrier and causes sedation.

**DELIRIOUS:**
- Consider Haloperidol 500 micrograms SC 2 hourly PRN or Levomepromazine 6.25mg SC 4-6 hourly PRN (monitor for extrapyramidal side effects)
  - If 2 or more doses of medication required to settle the patient in a 24hour period consider setting up a continuous subcutaneous infusion (CSI)

**Agitation**
- **Dose Range via CSCI/24 hours**
  - Midazolam for agitation: up to 30mg
  - Haloperidol for agitation: up to 5mg
  - Levomepromazine: up to 25mg
  - Seek Specialist Palliative Care advice if doses above 30mg of Midazolam, 5mg of Haloperidol or 25mg of Levomepromazine are needed

## QUICK GUIDE AGITATION / TERMINAL RESTLESSNESS IN THE LAST DAYS OF LIFE

## Reference
Merseyside and Cheshire Palliative Care Network Audit Group Guidelines on Symptom Control in the Dying Person

## Assessment/Description
Look for any reversible cause of agitation, and if identified institute appropriate management plans, such as inserting a urinary catheter for urine retention, constipation, full stomach, managing pain, etc. Consider and where possible address physical, psychological and spiritual factors as well as environmental factors such as light and noise.

**Pharmacological Options:**

**INITIALLY:**
- Midazolam 2.5mgs - 5 mgs SC up to 2 hourly PRN.
  - If eGFR < 30 give a reduced dose of Midazolam e.g. 1 mg to 2.5 mgs SC PRN

**ONGOING:**
- Consider Haloperidol 500 micrograms SC 2 hourly PRN or Levomepromazine 6.25mg SC 4-6 hourly PRN (monitor for extrapyramidal side effects)
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## QUICK GUIDE NAUSEA AND VOMITING IN THE LAST DAYS OF LIFE

**Reference**
Merseyside and Cheshire Palliative Care Network Audit Group Guidelines on Symptom Control in the Dying Person

### Assessment/Description
Patient complains of nausea, or is vomiting

### Non-Pharmacological options
- Reposition patient
- Eliminate known precipitants / strong odours
- Acknowledgement and explanation

### Pharmacological Options:

**INITIALLY:**
- Levomepromazine 2.5 - 6.25mg SC 6 hourly PRN (max dose 25mg / 24hour)
  - Lower dose may avoid undue sedation in some patients
  - See below for alternative anti-emetics

**ONGOING:**
- Continue to use Levomepromazine 2.5 - 6.25mg SC PRN 6 hourly
  - Review dosage after 24 hours.
  - If 2 or more doses given consider a CSCI with 6.25-12.5mg over 24 hour

Alternative anti-emetics include:
- Haloperidol 500 micrograms – 1.5mg SC PRN 8 hourly (max dose 5mg/ 24hour)
- Cyclizine 50mg SC PRN 8 hourly (max dose 150mg / 24hour)

Nausea and vomiting can be complex to manage - if patient is not settling seek Specialist advice

Remember that medication can be a profound cause of nausea and vomiting as can psychological issues

Raised intracranial pressure due to brain metastases may cause nausea and/or vomiting that might respond to high dose steroids (3.3mgs - 6.6mgs Dexamethasone SC OD)

See page 10 for further information

## QUICK GUIDE BREATHLESSNESS IN THE LAST DAYS OF LIFE

**Reference**
Merseyside and Cheshire Palliative Care Network Audit Group Guidelines on Symptom Control in the Dying Person

### Assessment/Description
Breathlessness causes significant anxiety. If heart failure is a contributing factor consider a trial of a diuretic via a suitable route.

Only use oxygen if patient has been shown to be hypoxic, the aim is for comfort not to maintain oxygen saturations.

Do not exceed oxygen flow rates of 4 litres/min except with Specialist advice.

### Non-Pharmacological options
- Reposition patient - Sit up / lean forward
- Acknowledgement and explanation
- Gentle air flow with fan / open window
- Regular mouth care

### Pharmacological Options:

**INITIALLY:**
- If patient not on an opioid regularly:
  - Morphine 2.5 - 5mg SC 4 hourly PRN. Or 2.5 - 5mg PO 4 hourly PRN if safe swallow

- If patient on an opioid regularly use:
  - Midazolam 2.5- 5mg SC PRN 2 hourly

**ONGOING:**
- If 1-2 PRN doses required in 24 hours set up a CSCI with either:
  - Morphine 10mg /24hour via CSCI or
  - Midazolam 10mg / 24hour via CSCI
  - (Midazolam is a useful option in a patient with renal failure if there is doubt about opioid choice)

**ONGOING:**
- Continue with rescue doses of either
  - Morphine 2.5 - 5mg SC PRN 4 hourly
  - Midazolam 2.5 - 5mg SC PRN 2 hourly
  - If not settling, seek Specialist advice
SIGNIFICANT RENAL IMPAIRMENT - SEEK SPECIALIST PALLIATIVE CARE ADVICE

- Paracetamol at standard doses is safe in renal impairment.
- If the eGFR is below 30mls/min (CKD 4/5) there is an increased risk of toxic side effects with all opioids due to drug and metabolite accumulation. Opioids should therefore be used with caution and should be monitored on a regular basis. Watch for signs of opioid toxicity which may include hallucinations, myoclonic jerks, drowsiness or confusion.
- When prescribing oral (strong) opioids, the immediate release forms are preferred. Long-acting opioid preparations should be avoided (e.g. MST/MXL) as the metabolites accumulate in renal failure. Fentanyl patches may be better tolerated in significant renal impairment but are difficult to titrate if pain is rapidly changing.
- Whilst parenteral Alfentanil or Fentanyl are pharmacokinetically the safest analgesics to use in renal failure as the metabolites are non-toxic, they may not be available in all localities and Oxycodone or Morphine or Diamorphine at reduced doses and / or frequency may be used but seek Specialist Palliative Care advice.
- NSAIDS should be avoided if possible, unless a patient is already on dialysis. If an NSAID must be prescribed for clinical reasons, the lowest effective dose should be used and the renal function should be re-checked within 5-7 days of starting the drug. If the renal function deteriorates further then a clinical decision is needed as to the benefits of continuing it's use.
- Adjuvant analgesics: Gabapentin / Pregabalin are safe in mild renal failure but if eGFR is less than 20ml/min the dose and / or frequency may need to be reduced to avoid toxicity.
- Anti-emetics: Haloperidol is the drug of choice for nausea in patients with renal failure, but if eGFR is less than 10ml/min the dose should be reduced (250 micrograms to 500 micrograms PO or SC). Levomepromazine is an alternative starting at 3mg PO or 2.5mgs SC. Adjust dose depending on effectiveness and side effects. Cyclizine should be avoided due to the risk of hypotension / tachyarrhythmia. Metoclopramide should be avoided due to the increased risk of extrapyramidal reactions.

ALWAYS Seek Specialist advice from palliative care and the patient's renal unit for patients managed with Haemodialysis or Peritoneal Dialysis

CLINICALLY ASSISTED HYDRATION (CAH) AT THE END OF LIFE

Nutrition and hydration are often emotional topics for families and patients when approaching end of life. There is need for ongoing sensitive discussions about goals of care and realistic expectations of treatment. The views of the patient and any Advance Care Planning should be considered throughout, and support for the carers when these decisions are being made is essential.

Within palliative care, clinically assisted hydration, either via intravenous or subcutaneous (SC) infusion, is provided with the intent of improving quality of life. SC fluids involve less discomfort, have fewer potential adverse effects than the intravenous (IV) route, may be provided in multiple care settings and are cheaper to provide. SC fluids should not be used to resolve severe dehydration, in emergency situations, or in patients with fluid overload.

There may be practical difficulties when considering SC fluids in the community setting. Equipment and training may be required. Refer to local guidelines and policy.

Due to the lack of any clear evidence, decisions to initiate Clinically Assisted Hydration will vary from patient to patient depending on the estimated burden to benefit balance. Treatment should always be in conjunction with other quality care, including good mouth care.

Potential indications
- Symptomatic dehydration
- Thirst (may be unrelated to fluid status)
- Reversible renal impairment
- Opioid toxicity
- Excess sedation
- Family/patient distress.

Potential complications
- Line discomfort/infection
- Oedema/ascites/effusions
- Worsening secretions
- Increased symptom burden as a result of above
- Family/patient distress
- Systemic fluid overload.

Management

There should be an agreed, clear indication of what is to be achieved by administering CAH, which should be discussed with the patient and family. Isotonic or hypotonic solutions only should be used (e.g. 0.9% NaCl). Rate of infusion will vary by patient, but is generally gravity fed with around 1 litre of fluid administered per 24hours. Infusion site should be under regular review for signs of infection, fluid accumulation or discomfort (at least every 48 hours).
QUICK GUIDE  CORTICOSTEROIDS IN PALLIATIVE CARE (Follow local guidelines if available)

Reference  Northern Strategic Clinical Network’s Guidelines

Corticosteroids are used extensively in palliative care. Dexamethasone is the preferred choice due to its relatively high anti-inflammatory potency and lower incidence of fluid retention and biochemical disturbance. Whilst highly effective they should be used with caution and be constantly monitored to prevent avoidable complications. (Potency: Dexamethasone 1mg ~ Prednisolone 7.5mg).

Dexamethasone should be prescribed in terms of the ‘base’ (Dexamethasone) rather than the ‘salt’ (Dex Phosphate or Dex Sodium Phosphate). Tablets are formulated as the base. Prescribing injections can appear confusing. For practical purposes: 3.3mg/ml injection may be considered equal to 4mg tablet.


Treatment and Management

Standard starting doses for the different indications are not well established and must take account of patient factors. Ensure daily dose is administered before noon in order to minimise insomnia. Clinical response must be reviewed within 7 days. Titrate down to minimum effective dose as soon as is possible.

Anorexia: 2 - 6mg daily. Judge response within 2 weeks. Although enhanced effect can still be present at 4 weeks, short courses are recommended to reduce risk of side effects.

Adjuvant analgesic: 8 - 16mg in cancer-related pain (e.g. liver capsular pain, nerve compression).

Anti-emetic: for chemotherapy follow Oncology guidelines. Refractory nausea and vomiting: 8 - 16mg daily.

Obstructive syndromes e.g. bowel obstruction, upper airways compression, SVCO, lymphangitis carcinomatosis: 6 - 16mg daily.

Spinal cord compression: 16mg daily for 5 days. Maintain on 8mg daily during radiotherapy, then reduce dose over 2 weeks.

If symptoms recur, increase to previous effective dose for at least 2 weeks before reducing again.

Raised intracranial pressure: 8 - 16mg daily for one week, and then reduce over 2-4 weeks to lowest dose which maintains benefit. (If treated with radiotherapy, steroids should be continued until one week post treatment, and then reduced as above). Consider trial of dose increase if symptoms recur.

ADVERSE EFFECTS:

- Glucose metabolism: Steroids can increase blood sugar levels. All patients on steroids should have regular blood glucose checks as per local guidance
- Insomnia: Give single or divided daily dose before noon to prevent insomnia.
- Dyspepsia: Give after food. Co-prescribe PPI if history of peptic ulcer disease or patient also taking Aspirin, NSAIDs, SSRIs or is anti-coagulated with Warfarin, LMWH or other agent.
- Psychiatric disturbance: depression, mania, psychosis, delirium.
- Change in appearance: moon face, truncal obesity, negative body image.
- Musculoskeletal problems: proximal myopathy, osteoporosis, avascular bone necrosis.
- Increased susceptibility to infection: especially oral/pharyngeal candidosis (examine mouth regularly).
- Skin changes: thinning, bruising, acne, impaired wound healing.
- Other: hypertension, oedema, pancreatitis.

SAFE USE: Monitoring and stopping treatment

Use the lowest effective dose for the shortest period of time. Close careful monitoring is essential. Steroid withdrawal: stop without tapering dose if total treatment duration of less than 3 weeks AND daily Dexamethasone dose of 6mg or less AND symptoms unlikely to relapse.

Gradual dose reduction: is necessary if any of following:

- 3 or more weeks treatment, daily dose of more than 6mg Dexamethasone,
- Risk of recurrent severe symptoms,
- Repeated courses of steroids,
- Other possible causes of adrenal suppression.

Daily dose can be reduced rapidly (e.g. halving dose) to 4mg/day, then more slowly by 1 - 2mg weekly in order to prevent a hypoadrenal crisis (malaise, profound weakness, hypotension).

Steroid treatment card: Patients on systemic steroids for > 3 weeks must be given a steroid card.

STEROIDS in last days of life: For ongoing symptom control, continue at the most convenient SC dose. If recent and/or low oral dose prescription for appetite stimulation, discontinue. If long-term, oral dose for whatever indication consider continuing at physiological dose, Dexamethasone 1.1mg SC.
Proactive Identification Guidance – proactively identifying patients earlier.

This updated 6th edition of the GSF PIG, renamed as Proactive Identification Guidance and formally known as Prognostic Indicator Guidance, aims to enable the earlier identification of people nearing the end of their life who may need additional supportive care. This includes people who are nearing the end of their life following the three main trajectories of illness for expected deaths – rapid predictable decline e.g. cancer, erratic decline e.g. organ failure and gradual decline e.g. frailty and dementia. Additional contributing factors when considering prediction of likely needs include current mental health, co-morbidities and social care provision.

Three trajectories of illness (Lynn et al) reflecting the three main causes of expected death

1. Rapid predictable decline e.g. Cancer
2. Erratic unpredictable e.g. Organ Failure
3. Gradual decline e.g. Frailty, Dementia, multi-morbidity

Why is it important to identify patients early?

Earlier recognition of decline leads to earlier anticipation of likely needs, better planning, fewer crisis hospital admissions and care tailored to peoples’ wishes. This in turn results in better outcomes with more people living and dying in the place and manner of their choice. Once identified, people are included on a register and where available the locality/electronic register, triggering specific active supportive care, as used in all GSF programmes and in GSF cross boundary care sites.

The 3 key steps of GSF

PIG and GSF – Early proactive identification of patients is the crucial first step of GSF, used by many thousands of doctors and nurses in the community and hospitals. For more information on GSF, how it is used in practice to help identify patients early, assess needs and wishes through advance care planning discussions and plan care tailored to patient choices, see the GSF website.

National Policy support for earlier identification.

General Medical Council – 2010

www.gmc-uk.org/static/documents/content/End_of_life.pdf

The GMC definition of End of Life Care; ‘People are ‘approaching the end of life’ when they are likely to die within the next 12 months. This includes people whose death is imminent (expected within a few hours or days) and those who:

- Advanced, progressive, incurable conditions.
- General frailty and co-existing conditions that mean they are expected to die within 12 months.
- Existing conditions if they are at risk of dying from a sudden acute crisis in their condition.
- Life threatening acute conditions caused by sudden catastrophic events.’

NICE Guidance in End of life care 2011 Quality statement 1

https://www.nice.org.uk/guidance/qs13/chapter/Quality-statement-1-Identification

- ‘Identification – People approaching the end of life are identified in a timely way.
- Systems – Evidence of local systems in place to document identification of people approaching the end of life.’

Proactive Identification Guidance – GSF PIG Flow-chart
For patients with advanced disease or progressive life limiting conditions, would you be surprised if the patient were to die in the next year, months, weeks, days? The answer to this question should be an intuitive one, pulling together a range of clinical, social and other factors that give a whole picture of deterioration. If you would not be surprised, then what measures might be taken to improve the patient’s quality of life now and in preparation for possible future decline?

General physical decline, increasing dependence and need for support.
- Repeated unplanned hospital admissions.
- Advanced disease – unstable, deteriorating, complex symptom burden.
- Decreasing activity – functional performance status declining (e.g. Barthel score) limited self-care, in bed or chair 50% of day and increasing dependence in most activities of daily living.
- Decreasing response to treatments, decreasing reversibility.
- Patient choice for no further active treatment and focus on quality of life.
- Progressive weight loss (>10%) in past six months.
- Sentinel Event e.g. serious fall, bereavement, transfer to nursing home.
- Serum albumin <25g/l.
- Considered eligible for DS1500 payment.

At least two of the indicators below:
- Patient for whom the surprise question is applicable.
- CHF NYHA Stage 3 or 4 with ongoing symptoms despite optimal HF therapy – shortness of breath at rest on minimal exertion.
- Repeated admissions with heart failure – 3 admissions in 6 months or a single admission aged over 75 (50% 1yr mortality).
- Difficult ongoing physical or psychological symptoms despite optimal tolerated therapy.
- Additional features include hyponatraemia <135mmol/l, high BP, declining renal function, anaemia, etc.

Liver Disease
Hepatocellular carcinoma.
Liver transplanted contra indicated.
Advanced cirrhosis with complications including:
- Refractory ascites
- Encephalopathy
- Other adverse factors including malnutrition, severe comorbidities, Hepatorenal syndrome
- Bacterial infection current bleeds, raised INR, hyponatraemia, unless they are a candidate for liver transplantation or amenable to treatment of underlying condition.

General Neurological Diseases
- Progressive deterioration in physical and/or cognitive function despite optimal therapy.
- Symptoms which are complex and too difficult to control.
- Swallowing problems (dysphagia) leading to recurrent aspiration pneumonia, sepsis, breathlessness or respiratory failure.
- Speech problems: increasing difficulty in communications and progressive dysphasia.

Parkinson’s Disease
- Drug treatment less effective or increasingly complex regime of drug treatments.
- Reduced independence, needs ADL help.
- The condition is less well controlled with increasing “off” periods.
- Dyskinesias, mobility problems and falls.
- Psychiatric signs (depression, anxiety, hallucinations, psychosis).
- Similar pattern to frailty – see below.

Motor Neurone Disease
- Marked rapid decline in physical status.
- First episode of aspirational pneumonia.
- Increased cognitive difficulties.
- Weight Loss.
- Significant complex symptoms and medical complications.
- Low vital capacity (below 70% predicted spirometry), or initiation of NIV.
- Mobility problems and falls.
- Communication difficulties.

Multiple Sclerosis
- Significant complex symptoms and medical complications.
- Dysphagia – poor nutritional status.
- Communication difficulties e.g., Dysarthria – fatigue.
- Cognitive impairment notably the onset of dementia.

3. Frailty, dementia, multi-morbidity

Frailty
For older people with complexity and multiple comorbidities, the surprise question must triangulate with a tier of indicators, e.g. through Comprehensive Geriatric Assessment (CGA).
- Multiple morbidities.
- Deteriorating performance score.
- Weakness, weight loss exhaustion.
- Slow Walking Speed – takes more than 5 seconds to walk 4 m.
- TUG – time to stand up from chair, walk 3 m, turn and walk back.
- PRISMA – at least 3 of the following:
  - Aged over 85, Male, Any health problems that limit activity, Do you need someone to help you on a regular basis?, Do you have health problems that cause require you to stay at home?, In case of need can you count on someone close to you?, Do you regularly use a stick, walker or wheelchair to get about?

Dementia
Identification of moderate/severe stage dementia using a validated staging tool e.g., Functional Assessment Staging has utility in identifying the final year of life in dementia. (BGS) Triggers to consider that indicate that someone is entering a later stage are
- Unable to walk without assistance and
- Urinary and faecal incontinence, and
- No consistently meaningful conversation and
- Unable to do Activities of Daily Living (ADL)
- Barthel score >3
Plus any of the following: Weight loss, Urinary tract Infection, Severe pressures sores – stage three or four, Recurrent fever, Reduced oral intake, Aspiration pneumonia. NB Advance Care Planning discussions should be started early at diagnosis.

Stroke
- Use of validated scale such as NIHSS recommended.
- Persistent vegetative, minimal conscious state or dense paralysis.
- Medical complications, or lack of improvement within 3 months of onset.
- Cognitive impairment / Post-stroke dementia.
- Other factors e.g. old age, male, heart disease, stroke sub-type, hyperglycaemia, dementia, renal failure.

The Surprise Question
- Evidence of local systems in place to document
- Regularly reassess
- NO
- Reassess
- YES

Liver Disease continued
- Advanced disease – unstable, deteriorating, complex symptom burden.
- Presence of significant multi-morbidities.
- Decreasing activity – functional performance status declining (e.g. Barthel score) limited self-care, in bed or chair 50% of day and increasing dependence in most activities of daily living.
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