The Palliative Care Handbook

guidelines for clinical management
and symptom control

Rod MacLeod, Jane Vella-Brincat, Sandy Macleod
THE PALLIATIVE CARE HANDBOOK

GUIDELINES FOR CLINICAL MANAGEMENT AND SYMPTOM CONTROL

8TH EDITION 2016

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Many of the medications listed are being used outside their product licence. Prescription of a drug (whether licensed use/route or not) requires the prescriber, in the light of published evidence, to balance both the potential good and the potential harm which might ensue. Prescribers have a duty to act with reasonable care and skill in a manner consistent with the practice of professional colleagues of similar standing. Thus, when prescribing outside the terms of the licence, prescribers must be fully informed about the actions and uses of the drug, and be assured of the quality of the particular product (www.palliativedrugs.com/using-licensed-drugs-for-unlicensed-purposes). Prescribers also have a duty to inform patients that drugs are being used outside their licence and to inform them of any expected effects and side effects. Care has been taken to ensure accuracy of information at time of printing. This information may change and final responsibility lies with the prescriber. Some medication will incur a cost to the user, it is important to consider this before prescribing.

This Handbook should be used in conjunction with Therapeutic Guidelines – Palliative Care – version 3 (Therapeutic Guidelines Limited, Melbourne) where possible

Throughout the book, drugs that are either not available or not funded in New Zealand are marked with *

Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>subcut</td>
<td>subcutaneous</td>
</tr>
<tr>
<td>CNS</td>
<td>central nervous system</td>
</tr>
<tr>
<td>bd</td>
<td>twice daily</td>
</tr>
<tr>
<td>LFTs</td>
<td>liver function tests</td>
</tr>
<tr>
<td>tds</td>
<td>three times daily</td>
</tr>
<tr>
<td>MAOIs</td>
<td>monoamine oxidase inhibitors</td>
</tr>
<tr>
<td>qid</td>
<td>four times daily</td>
</tr>
<tr>
<td>NSAIDs</td>
<td>nonsteroidal anti-inflammatory drugs</td>
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Printed by

Crucial Colour

FOREWORD

Palliative care has come a long way since the inception of the modern hospice movement in the UK in the 1960s. It is now recognised as an integral and important part of health care in over 100 countries throughout the world. In New Zealand, the growth of the specialty has been significant and now people throughout New Zealand should expect that those who care for people at the end of life should have access to the best quality of care.

This handbook – which started life as a small pocket book written by Rod MacLeod and Jane Vella-Brincat in Bath, England in 1994 has now grown to become an invaluable resource for practitioners throughout the country. Its popularity stems from the ease of use, the basic layout and the uncomplicated explanations of how to manage challenging symptoms that the authors have developed.

This handbook finds its place in all areas where palliative care happens; it gives confidence to those who use it and hopefully therefore comfort to those approaching death so that they live every moment.

Mary Schumacher
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Wellington, NZ
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INTRODUCTION

The first section of this book is a set of guidelines for the alleviation of symptoms commonly encountered in palliative care. Drug therapy is included.

The second section (the pharmacopoeia) contains drug information:

- it is in alphabetical order by generic drug name
- the interactions listed include discussion about enzymes responsible for drug metabolism commonly known as Cytochrome P450 (CYP) enzymes. There are many CYP enzymes some of which are genetically controlled. The interactions listed are based mainly on theory, are subject to change as more is learnt about the CYP enzyme system and are meant to be used as a guide only to potential interactions. Only commonly used palliative care drugs have been included but interactions with other drugs may also occur.
- there is also information about the use of syringe drivers
PALLIATIVE CARE AIMS

- to achieve the best possible quality of life for patients and their families
- to understand and address patients’ physical, psychological, social and spiritual suffering
- to be applicable from early on in the course of the illness

THE WORLD HEALTH ORGANISATION DEFINED PALLIATIVE CARE AS:
‘An approach that improves the quality of life of patients and their families facing the problems associated with a life-threatening illness, through the prevention and relief of suffering by means of early identification and impeccable assessment and treatment of pain and other problems, physical, psychosocial and spiritual.’

Palliative care:
- provides relief from pain and other distressing symptoms
- affirms life and regards dying as a normal process
- intends neither to hasten or postpone death
- integrates the psychological and spiritual aspects of patient care
- provides support to help patients live as actively as possible
- provides support to the family during the illness and bereavement
- uses a multidisciplinary team approach
- enhances quality of life and influences the course of the illness
- is applicable early in the course of illness alongside therapies that are intended to prolong life (e.g. chemotherapy, radiotherapy) and diagnostic investigations

GENERAL SYMPTOM MANAGEMENT PRINCIPLES

- accurate and meticulous assessment is essential
- assess and address non-physical as well as physical issues
- difficult to control symptoms may require several different approaches
- aim for highest possible quality of life
- use risk versus benefit assessments when side effects of therapy occur
- explain issues as much as possible to the patient and their carers
- use a multidisciplinary approach
- reassess continuously
PAIN

The assessment and management of pain and other symptoms are the cornerstones of effective palliative care. There are different types of pain and many patients have more than one.

COMPREHENSIVE ASSESSMENT

• listen to the patient’s story and the language used
• ask about the site(s) of pain
• measure intensity with a validated tool to assess changes:
  — a visual analogue scale (some patients find this hard to use)
  — a numerical rating scale – perhaps the most common method used – patients rate their pain on a scale of 0 (no pain) to 10 (the worst pain they can imagine)
  — colour charts
  — facial expression charts
• ask about timing and duration of pain e.g. constant or episodic
• ask about the nature (e.g. stabbing, aching) and duration of the pain – this will determine management
  — identifies the type and source of pain
    > somatic nociceptive is usually constant and localised
    > visceral is usually described as deep or aching (capsular stretch pain) or intermittent and griping (colicky pain)
    > bone pain is usually deep or boring
    > neuropathic pain is usually burning, shooting or stabbing
• ask about what relieves the pain (body position, heat, cold) and what exacerbates the pain (movement, position, heat)
• ask about the significance of the pain
  — ask how much of a nuisance it is
  — discuss its significance
  — explain the likely causes – often helpful in allaying fears or anxieties and can significantly contribute to the relief of pain
• examine the part(s) that are painful – look, touch and move
• consider further investigation such as X-ray, CT or MRI but only if the result is going to influence management
• document all findings to compare and communicate
• review regularly – essential after any therapeutic intervention

OTHER ASSESSMENT FACTORS

In a bio-medical model of practice it is tempting to assume that pain has a predominant physical component. Often, physical pain is only part of the symptom complex (through direct or indirect tumour effects or non-malignant processes). Psychological, spiritual and sociological elements will also be identifiable in many people with pain. Fear, anxiety, sadness, anger, frustration and isolation are but
a few of the feelings that can contribute to the total perception of pain. All of these elements help to build a realistic picture of the overall impact of pain on the individual’s quality of life.

MANAGEMENT

It is important to encourage patients to develop self-management strategies and to utilise non-pharmacological strategies such as rest, positioning, pacing etc. There are also a number of enabling strategies like goal setting, pain management plans, scripts and diaries that many will find useful.

ANALGESICS

- some pains may not respond completely to opioids
- co-analgesics are useful when response to opioids is poor
- switching route can sometimes help e.g. from oral to subcutaneous
- in prescribing analgesics use a step-wise approach:

<table>
<thead>
<tr>
<th>Analgesics</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>morphine</td>
<td>or oxycodone</td>
</tr>
<tr>
<td>or hydromorphone*</td>
<td>or fentanyl</td>
</tr>
<tr>
<td>or methadone</td>
<td></td>
</tr>
<tr>
<td>codeine</td>
<td>or dihydrocodeine</td>
</tr>
<tr>
<td>or tramadol</td>
<td>or buprenorphine</td>
</tr>
<tr>
<td>regular paracetamol</td>
<td></td>
</tr>
</tbody>
</table>

- regular paracetamol may be useful in opioid induced hyperalgesia although use should be continued only if effective as up to 8 tablets per day adds significantly to the tablet burden
- there is some debate over the second step in this ladder
  - most palliative care practitioners go to step 3 either after step 1 or initially depending on the severity of the pain
  - pain relief from codeine may be from the active metabolite, morphine
  - the place of tramadol in palliative care remains unclear – it can be extremely emetogenic

Initiating morphine in opioid naïve patients

- start with small regular oral (if possible) immediate release doses
- titration with slow release morphine is less common than with immediate release
- if using immediate release prescribe morphine elixir (immediate release) (2.5 to 5 mg) every four hours regularly and titrate
- prescribe ‘when required’ doses of 1/5th to 1/6th of the regular 24 hour dose for ‘breakthrough’, ‘episodic’ or ‘incident’ pain
• document the amount of morphine taken
• once a stable dosing regimen is achieved (2 to 3 days) convert to a long-acting preparation
  — calculate the total 24 hour dose of immediate release morphine required from ‘breakthrough’ and regular dosing, divide by two and give twice daily
  — ‘when required’ doses of 1/5th to 1/6th of the regular 24 hour dose should be prescribed as immediate release once again for pain between doses
• if the patient can no longer swallow
  — give ½ the total 24 hour oral dose by continuous subcutaneous infusion
  — ‘when required’ doses of 1/5th to 1/6th of the regular 24 hour dose should be prescribed once again for pain between doses
• consider reducing dose if another mode of pain relief is used (e.g. radiotherapy, ketamine)

Initiating oxycodone in opioid naïve patients
• start with small regular oral (if possible) doses
• prescribe oxycodone immediate release capsules or liquid every 4 to 6 hours and titrate
• prescribe ‘when required’ doses of 1/10th to 1/12th initially (although many practitioners use 1/5th to 1/6th) of the regular 24 hour dose for ‘breakthrough’, ‘episodic’ or ‘incident’ pain
• document the amount of oxycodone taken
• once a stable dosing regimen is achieved (2 to 3 days) convert to a long-acting preparation
  — calculate the total daily dose of oxycodone required from ‘breakthrough’ and regular dosing, divide by two and give twice daily
  — ‘when required’ doses of 1/10th to 1/12th initially (although many practitioners use 1/5th to 1/6th) of the regular 24 hour dose should be prescribed as immediate release for pain between doses
• consider reducing dose if another mode of pain relief is used (e.g. radiotherapy, ketamine)
• the long acting preparation has a layer of immediate acting drug round it
• if the patient can no longer swallow
  — give ½ the total 24 hour oral dose by continuous subcutaneous infusion
  — ‘when required’ doses of 1/10th to 1/12th initially (although many practitioners use 1/5th to 1/6th) of the regular 24 hour dose should be prescribed once again for pain between doses
• consider reducing dose if another mode of pain relief is used (e.g. radiotherapy, ketamine)

Initiating hydromorphone* in opioid naïve patients
• as with oxycodone start with small regular oral (if possible) doses
• titrate upwards in small increments as with morphine and oxycodone
• otherwise the same principles apply
Initiating fentanyl patches in opioid naïve patients
• don’t – fentanyl patches should only be used in patients who have already been exposed to opioids

Initiating methadone in opioid naïve patients
• as methadone has a long and variable half life it should be commenced at low dosage e.g. 1 mg to 2.5 mg bd and consideration should be given to dose reduction once at steady state (minimum 5 days)
• should be used under advice of a specialist palliative care physician only

Adverse effects of opioids
• all opioids are associated with the following adverse effects but the incidence (incidences below are for morphine) and severity vary from opioid to opioid (e.g. fentanyl is less constipating than morphine)
• tolerance to some of these adverse effects can develop e.g. nausea/ vomiting but not to others e.g. constipation
  — constipation – 95% of patients (less with fentanyl [50%] and the naloxone/ oxycodone combination product) – prescribe a laxative prophylactically
  — nausea/vomiting – 30-50% of patients – usually in the first 10 days until tolerance develops
  — drowsiness – 20% of patients – usually in the first 3 to 5 days until tolerance develops
  — confusion – 2% of patients – either reduce the dose, change to a different opioid or consider adding haloperidol
  — hallucinations/nightmares – 1% of patients – give haloperidol or change to a different opioid
  — hyperalgesia – usually to touch as a result of too high a dose of opioid which may improve on dose reduction
  — hyperkatafèia – emotional lability induced by long-term opioid use

Opioid rotation
• opioid rotation (or changing from one opioid to another) is often used when tolerance to the analgesic effects of opioids (stimulation of NMDA receptors) or severe adverse effects occur
• works because of the difference in the mix of opioid receptors stimulated by each individual opioid in each patient
• most often from morphine to oxycodone, fentanyl or methadone
• rotation should only occur under supervision and by a specialist as conversion doses are difficult to predict and are often much smaller doses than those listed below – see oxycodone, hydromorphone*, fentanyl and methadone in the second section
Opioid equivalents

- the following are ‘single dose’ equivalences i.e. ONLY equivalents in healthy volunteers given a single dose
- equivalence in sick patients who are chronically dosed is difficult to quantify – use care when converting from one opioid to another
- pethidine is NOT recommended in palliative care

<table>
<thead>
<tr>
<th>Drug</th>
<th>Oral Dose (mg)</th>
<th>Subcut Dose (mg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Codeine</td>
<td>60</td>
<td>5</td>
</tr>
<tr>
<td>Tramadol</td>
<td>100</td>
<td>2</td>
</tr>
<tr>
<td>Oxycodone</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>Methadone</td>
<td>see methadone</td>
<td></td>
</tr>
<tr>
<td>Fentanyl</td>
<td>see fentanyl</td>
<td></td>
</tr>
<tr>
<td>Buprenorphine</td>
<td>see buprenorphine</td>
<td></td>
</tr>
</tbody>
</table>

Drugs that are either not available or not funded in New Zealand are marked with *

Co-analgesics

- drugs usually used for a different indication with analgesic properties (sometimes such use is outside the product license)
- can be used in combination with other analgesics or alone
- choice is determined by the types of pain
- the use of co-analgesics is probably most helpful in neuropathic pain

- **bone pain** – due to tumour or metastatic involvement
  - NSAIDs e.g. diclofenac – inhibit prostaglandins
  - bisphosphonates e.g. pamidronate, zoledronic acid
  - denosumab*
- **skeletal muscle spasm pain** – due to tumour involvement
  - muscle relaxants e.g. diazepam, clonazepam, baclofen
- **smooth (intestinal) muscle spasm pain** – ‘colic’ from intestinal spasm
  - anticholinergic/antimuscurinic e.g. hyoscine butylbromide
- **tenesmus** – due to tumour or metastatic involvement of the rectal muscles
  - steroids e.g. dexamethasone, prednisone – decrease inflammation around tumour
- **raised intracranial pressure** – due to tumour or fluid
  - steroids e.g. dexamethasone – decrease inflammation around tumour
  - NSAIDs e.g. diclofenac – inhibit prostaglandins
- **liver capsule stretch pain** – from an enlarged liver
  - steroids e.g. dexamethasone – decrease inflammation
NEUROPATHIC PAIN
• often the most severe and difficult to manage of all persisting pains
• caused by damage to the nervous system
• involves NMDA receptor stimulation to some extent
• severity cannot usually be linked to the amount of damage
  — ‘trivial’ lesions can produce severe pain

Causes
• peripheral nerve damage – post-surgical, post-trauma or compression
• herpetic nerve invasion
• amputation – phantom limb pain
• Chronic Regional Pain Syndrome (CRPS)
• nerve root injury – traumatic avulsion, post-spinal surgery
• epidural scarring, arachnoiditis
• spinal cord injury and disease
• stroke
• diabetes
• chemotherapy e.g vincristine, oxaliplatin, taxanes, cisplatin

Drugs that are either not available or not funded in New Zealand are marked with *

Characterisation
• characterised by description and by cause
  — BUT the pain is not always within the distribution of a dermatome or a peripheral nerve
• includes allodynia (pain in an area of altered sensitivity) and other sensory symptoms
• generally continual and of varying intensity
  — variability in intensity is spontaneous and often has a paroxysmal component not necessarily related to stimulation
• descriptive terms include burning, cutting, stabbing sharp/shooting
• crushing
• episodic pain, which can be present on top of the continuous pain, may itself be brief but often a long-lasting aching pain remains for several hours

Management
• a multidisciplinary approach is useful
• behavioural modification – any treatment will be of only limited value unless certain behaviours are changed so address cognitive, mood and behavioural aspects of the patient’s pain individually or in a group
• drugs
  — opioid analgesics (first line for neuropathic pain) should be trialed but doses may increase rapidly – some opioids may be more useful than others e.g. methadone which has NMDA blocking activity
  — centrally acting agents reduce spinal hyperexcitability
— some drugs have an effect on nociceptor neuromodulators, neurotransmitters and cell membrane stability
— efficacy is highly variable between drugs so tailor the drug to the patient
  > gabapentin, pregabalin*
  > anticonvulsants e.g. valproate
  > benzodiazepines e.g. clonazepam
  > tricyclic antidepressants e.g. nortriptyline SSRIs e.g. escitalopram, sertraline – limited efficacy in palliative care
  > SNRIs e.g.duloxetine*, venlafaxine
  > antiarrhythmics e.g. mexiletine
  > muscle relaxants e.g. baclofen
  > NMDA antagonists e.g. ketamine
  > alpha-adrenergic agents e.g. clonidine
  > calcium channel blockers e.g. nifedipine
  > steroids e.g. dexamethasone for nerve pressure pain
  > sodium channel blockers e.g. lignocaine
— combining an antidepressant with an anticonvulsant or similar may be more effective than either alone e.g. nortriptyline + gabapentin, pregabalin*
if the above are ineffective consider intrathecal/epidural opioids, local anaesthetics and clonidine

• other analgesic modalities
  — nerve blocks
    > availability is dependent on the skills of the team
    > access to a specialist anaesthetist is not always possible
    > for pain which breaks through analgesia, or is controlled at rest but not on movement or is nonresponsive
    > upper abdominal pain due to pancreatic cancers may respond to coeliac plexus blocks
  — others – often used in conjunction with analgesics
    > mobilisation e.g. structured stretching, progressive resistance training
    > radiotherapy/surgery
    > cytotoxic drugs
    > hormone therapy
    > spinal delivery systems
    > neuromodulation e.g. transcutaneous nerve stimulation (TENS) and, very occasionally, implanted devices such as peripheral nerve or spinal cord stimulation
NAUSEA/VOMITING

These are common symptoms in palliative care and are often difficult to control.

- it is important to separate nausea from vomiting
- consider how each affects the individual patient
  - a vomit a day with no nausea may be more acceptable than continuous low-level nausea
  - for some patients nausea is more distressing than pain
- nausea and/or vomiting often has more than one cause
- choose a management strategy to fit the cause(s)
- antiemetics work at differing sites and receptors
- antiemetics that affect multiple receptors in multiple areas, such as levomepromazine (methotrimeprazine), may be useful choices regardless of cause
- a combination of antiemetics is useful, particularly where there are multiple causes

Causes

There are two distinct areas in the central nervous system (CNS), which are predominantly involved with nausea and vomiting:

- chemoreceptor trigger zone (CTZ) close to the area postrema
  - part of the central nervous system, the CTZ is thought to lie outside the blood/brain barrier and so can be affected by causes and treatment which are unable to penetrate the CNS
- the vomiting centre in the medulla oblongata
  - can be directly stimulated or inhibited by certain agents

The CTZ sends impulses to the vomiting centre, which then initiates nausea and/or vomiting. Higher centres involved with fear and anxiety also communicate with the vomiting centre, as do the peripheral vagal and sympathetic afferents and the vestibular nerve.

The causes can be summarised as:

- higher centre stimulation – fear/anxiety
- direct vomiting centre stimulation – radiotherapy to the head, raised intracranial pressure
- vagal and sympathetic afferent stimulation – cough, bronchial secretions, hepatomegaly, gastric stasis, constipation, intestinal obstruction
- chemoreceptor trigger zone stimulation – uraemia, hypercalcaemia, drugs e.g. opioids, cytotoxics
- vestibular nerve stimulation – motion
Management

- higher centre stimulation (emotion – fear/anxiety)
  - counselling/explanation/listening
  - a benzodiazepine
- direct vomiting centre stimulation (radiotherapy to the head, raised intracranial pressure)
  - cyclizine
  - dexamethasone
- vagal and sympathetic afferent stimulation (cough, bronchial secretions, hepatomegaly, gastric stasis, constipation, intestinal obstruction)
  - cough – see cough
  - bronchial secretions – see retained secretions
  - constipation – see constipation
  - hepatomegaly
    - dexamethasone
    - cyclizine
  - gastric stasis
    - domperidone (minimal extrapyramidal effects)
    - metoclopramide
    - erythromycin – a strong prokinetic
  - intestinal obstruction
    - cyclizine
    - levomepromazine (methotrimeprazine)
    - avoid prokinetics e.g. metoclopramide in complete obstruction although use in partial obstruction may help – see intestinal obstruction section
- chemoreceptor trigger zone stimulation (uraemia, hypercalcaemia, drugs e.g. morphine)
  - haloperidol
  - levomepromazine (methotrimeprazine)
- vestibular nerve stimulation (motion)
  - cyclizine
  - hyoscine patch (scopolamine)
- other drugs which may be useful where others have failed
  - atypical antipsychotics e.g. olanzapine
  - ondansetron, palonosetron* (may cause constipation) – experience in palliative care is limited
  - aprepitant (a neurokinin 1 (NK1) antagonist from the class of drugs known as substance P antagonists) – used with steroids and ondansetron for delayed emesis following highly emetogenic chemotherapy. Its place in palliative care has not been established.
- other therapies with little evidence include acupuncture, ginger, cannabis

Drugs that are either not available or not funded in New Zealand are marked with *
BOWEL MANAGEMENT
• alteration in bowel function is common in terminally ill people
• constipation is more common than diarrhoea
• efficient bowel management may alleviate distress
• carefully assess bowel function on a daily basis
• regimens should be discussed, carried out and reported on daily

CONSTIPATION
• diagnose through an accurate history followed by examination
• it is the difficult or painful and infrequent passage of hard stools
• comparison with an individual’s normal bowel habit and usual use of laxatives may highlight changes related to disease or treatment
• a record of bowel habits will help in the management
• examination of the abdomen and the rectum may exclude faecal impaction or rectal pathology

Causes
• metabolic disturbances e.g. hypercalcaemia
• dehydration from vomiting, polyuria, sweating, tachypnoea
• drugs
  — cytotoxics e.g. vinca alkaloids (via neuropathies)
  — opioids via opioid receptors in the GI tract and perhaps in the CNS – > 95% of people taking morphine will become constipated although other opioids may be less constipating e.g. fentanyl, methadone
  — anti-cholinergics e.g. tricyclic antidepressants
  — aluminium salts in antacids
  — iron
  — antispasmodics e.g. hyoscine butylbromide
  — anti-Parkinsonian drugs e.g. levodopa
  — antipsychotics/anxiolytics
  — ondansetron, palonosetron*
• immobility e.g. weakness
• low fibre diet e.g. milky/invalid foods or reduced intake
• inability to obey the call to stool
• concurrent medical problems e.g. haemorrhoids, anal fissure, diabetes, hypothyroidism
• intestinal obstruction from tumour, faeces or adhesions (abdominal X-ray may help with diagnosis)
• gastrointestinal tract nerve compression or damage or autonomic neuropathy

Symptoms
• anorexia
• vomiting/nausea
• abdominal discomfort or cramping
• spurious diarrhoea or overflow
• confusion
• anxiety
• bowel obstruction
• pain

Management
• prevention is the key
• if a cause (or causes) are identified remove it (or them) if possible
• exercise reduces the risk of constipation so encourage it where possible
• encourage increased fibre e.g. bran, kiwi crush or soluble fibre formulations (require activity and fluids to avoid impaction)
• laxatives
  — when opioids are prescribed anticipate constipation and prescribe an oral softener with a stimulant laxative e.g. docusate with senna or bisacodyl which may prevent the need for rectal intervention later (NB if combinations cause cramps reduce the dose or use an osmotic laxative such as macrogol 3350 with electrolytes (Movicol™, Lax-Sachets™)
  — low dose opioid antagonists such as naloxone (marketed in combination with oxycodone and methylaltrexone*) are effective in opioid-induced constipation without affecting analgesia
  — if constipation is already present give a bisacodyl 10 mg suppository and a glycerin suppository or a sodium lauryl sulphoacetate enema (Micolette™)
  — avoid stimulant laxatives in people with signs of GI obstruction
  — if the patient has a partial obstruction use an osmotic/softener laxative e.g. docusate, and avoid stimulant laxatives
  — if the patient has a spinal cord compression where evacuation is difficult keep the bowel motion firm (avoid softeners) and use a stimulant
  — if a patient taking laxatives has no bowel motion for two days and this is not their normal bowel habit give extra laxatives and, if appropriate, kiwi fruit or prune juice
  — if a patient taking laxatives has no bowel motion for three days and this is not their normal bowel habit a rectal examination should be carried out
• if soft faeces are found give two bisacodyl 10 mg suppositories or one to two Micolette™ enemas
• if hard faeces are found give one or two glycerine suppositories or two bisacodyl 10 mg suppositories or consider macrogol 3350 with electrolytes (Movicol™, Lax-Sachets™)
• if rectum is empty (or no result from first action) repeat abdominal palpation and consider an abdominal X-ray
  — suppositories must make contact with the bowel wall to work
  — methylaltrexone*
• faeces consist of approximately 50% water, 25% bacteria and 25% food residue so even if the patient is not eating there will be faeces in the bowel
**DIARRHOEA**

- a relatively uncommon problem in palliative care
- rotation from morphine to fentanyl may result in a sudden reduction in opioid constipating effects resulting in diarrhoea

**Causes**

- faecal impaction (overflow) – identify with a clinical examination (including rectal)
- colo-rectal carcinoma (also causes discharge and tenesmus)
- loss of sphincter tone and sensation e.g. from spinal cord compression
- incomplete gastrointestinal obstruction – frequent or recurrent diarrhoea suggests partial obstruction so try lower bowel evacuation
- malabsorption or food intolerance e.g. from lack of pancreatic enzymes
- concurrent disease e.g. diabetes mellitus, hyperthyroidism, inflammatory bowel disease
- radiotherapy to the torso
- cytotoxics (e.g. capecitabine)
- antibiotics – C.difficile
- bowel surgery or inflammation
- anxiety
- opioid rotation to a less constipating opioid e.g. from morphine to fentanyl

**Management – dependent on cause**

- assess bowel habit and faecal consistency
- consider likelihood of infection
- maintain skin integrity around anal area – use barrier creams to prevent excoriation e.g. zinc oxide
- think about overflow from impaction or partial obstruction
- use abdominal examination or X-ray to rule out obstruction restrict oral intake (except fluids) to rest the bowel
- withhold laxatives where appropriate
- administer antidiarrhoeal medications such as loperamide, opioids
- if impacted use manual removal followed by laxatives
- in partial obstruction diarrhoea may be very unpleasant
- in spinal cord compression a constipating drug may help e.g. codeine, loperamide (although patients already receiving morphine may not benefit) followed by regular suppositories and/or manual removal
- in colo-rectal carcinoma a palliative colostomy or radiotherapy should be considered
- in malabsorption states, the addition of pancreatic enzymes at meal times will help the situation e.g. pancreatin or, in bile salt malabsorption, cholestyramine
- secretory diarrhoea (associated with carcinoid syndrome or AIDS) may respond to octreotide
- ondansetron or palonosetron* may be worth considering especially if nausea/vomiting are also present

Drugs that are either not available or not funded in New Zealand are marked with *
INTESTINAL OBSTRUCTION

Intestinal obstruction is a difficult area of palliative care. There is considerable inter-individual and intra-individual variation in symptoms and optimal management.

Causes
- can be mechanical or paralytic
- blockage of intestine by intraluminal or extraluminal tumour, inflammation or metastasis
- blockage can occur at multiple sites in patients with peritoneal involvement
- may be aggravated by drugs e.g. anticholinergics, opioids
- radiation fibrosis
- autonomic nerve disruption by tumour

Management
The management of intestinal obstruction should be tailored to the individual at the time with different strategies being employed when needed.
- explain the predicament
- give dietary advice e.g. foods with minimal residue
- minimise colic by stopping osmotic/stimulant laxatives (continue softeners) and give subcutaneous hyoscine butylbromide (20 mg bolus followed by 60 to 80 mg subcut infusion over 24 hours)
- give analgesia (commonly subcutaneous opioids)
- reduce vomiting by giving appropriate antiemetics e.g. cyclizine with or without haloperidol – metoclopramide should only be used if there is clear evidence that there is only a partial obstruction
- consider alternative measures e.g. surgery, radiotherapy
- steroids e.g. dexamethasone should be given a trial
- iv fluids and nasogastric tubes should be avoided but may be preferred where drug treatment has not worked. Subcut fluids may have a role in some
- somatostatin analogues (octreotide) may be used subcutaneously in specialist practice to reduce secretions and minimise symptoms
- if subacute intestinal obstruction, the aim may be to clear the obstruction using steroids e.g. dexamethasone to reduce the inflammation around the obstruction and hyoscine butylbromide to minimise secretions and colic then, at an appropriate time, to push gut contents through with a prokinetic agent e.g. metoclopramide
- the timings of each change in therapy will depend on the individual patient and their condition
- review the situation regularly
MOUTH CARE

Poor oral hygiene is probably the most significant factor in the development of oral disease near the end-of-life.

• good mouth care is essential to the well being of patients debilitated by advanced disease
• mouth problems are common – occurring in up to 90% of patients
• risk factors for oral problems include
  — debility, dry mouth (drugs, mouth breathing, radiotherapy), chemotherapy, dehydration, cachexia, weight loss, ill-fitting dentures

Assessment/causes

• appropriate and effective oral assessment should be carried out on each patient daily using a pen torch and spatula
• remember functions of saliva – cleansing and lubrication, buffering, remineralisation, antimicrobial, digestion, maintenance of mucosal integrity
• key questions for effective mouth care are
  — is the mouth dirty, dry, painful or infected?
  — also assess mental, nutritional and physical state, concurrent medications, tongue, teeth/dentures, mucous membranes, type of saliva, and lips
  — mental state will determine the patient’s ability and willingness to participate in their care
  — nutritional state will give an indication of the patient’s ability to chew and swallow as well as their general well being – a well balanced diet and adequate fluid intake are important in mouth care
  — physical state may also contribute to mouthcare issues e.g. low haemoglobin increases susceptibility to infections and may be accompanied by lethargy, weakness and dyspnoea, all of which contribute to mouth care problems
  — patients in pain may require extra help with their mouth care
  — concurrent medications can affect the state of the mouth e.g. opioids/antidepressants may cause dry mouth, steroids/antibiotics may encourage oral candidiasis
  — other causes of poor mouth care include debility, reduced oral intake, inability to brush teeth, dehydration, saliva-reducing drugs, chemotherapy or radiotherapy, oxygen therapy and mouth breathing

Management – prevention is a priority

• regular tooth and denture brushing, twice daily at least
• regular use of anti-bacterial and anti-fungal mouthwash
• check fit of dentures
• regular dental checks if possible
• regular mouthcare; frequency dictated by assessment
• check for infection
• check for bone or nerve damage
• check mucosa
• reduce caffeine and alcohol, diet drinks (have a low pH)
• hypersalivation may be helped with atropine eye drops 1%, 1 to 2 drops in the mouth three to four times a day, ipratropium bromide nasal spray, 1 to 2 puffs in the mouth three to four times a day, radiotherapy or botulinum toxin to salivary glands

**Dirty mouths**
• chlorhexidine mouthwash is a useful cleansing agent
• sodium bicarbonate mouthwash is used by many, especially in oncology
• there is little point in cleaning the mouth if dentures are worn unless those dentures are also meticulously cleaned (including soaking overnight in ¼ strength Milton™)

**Dry mouths**
• salivary stimulants e.g. lime juice, fresh melon or pineapple are useful in dry mouths as is a saliva substitute (often useful to freeze fruit first); also, lollies or mints (sorbitol, xylitol-containing gum)
• pilocarpine solution (1 mg/mL, 5 to 10 mL or 1-2 drops 4% eye drops rinsed three times a day) may be useful for dry mouths

**Infected mouths**
• nystatin suspension is useful in the treatment of oral candidiasis but may take up to two weeks to clear an infection and many candidal infections are now resistant to it
• miconazole oral gel is also useful in the treatment of oral candidiasis, usually after nystatin suspension has failed
• systemic anti-fungals e.g. fluconazole (50 mg a day for 7 to 14 days or 100-150 mg stat) are sometimes needed for intractable oral candidal infections
• aciclovir may be useful for herpetic infections

**Painful mouths**
• may need systemic opioids
• coating agents
  — sucralfate suspension (use crushed tablets)
  — topical anaesthesia e.g. lignocaine viscous (watch for choking hazards)
• benzydamine is an analgesic mouthwash for painful mouths
• topical corticosteroids e.g. triamcinolone in orabase may be useful for aphthous ulcers (not used if oral candidiasis present)
• Bonjela™ (choline salicylate) may soothe sore gums

**TASTE ALTERATION**
• reduction in taste sensitivity i.e. hypogeusia
• absence of taste sensation i.e. ageusia
• distortion of taste i.e. dysgeusia
Causes

- local disease of mouth and tongue
- systemic diseases
- partial glossectomy
- nerve damage
- zinc deficiency
- alteration to cell renewal via malnutrition, metabolic endocrine factors, viral infections, hyposalivation
- dental pathology/hygiene
- diabetes
- gastric reflux
- drugs
  - cyclizine
  - anticholinergics (leads to dry mouths)
  - chemotherapy
  - lithium
  - ACE inhibitors
  - citalopram (uncommon)

Management

- remove or treat causes e.g. give pilocarpine for dry mouth, stop likely drugs
- zinc (but only if zinc is deficient)
- use sialogogues such as chewing sugar-free gum or sour-tasting drops
- may be unresponsive to interventions

SWALLOWING DIFFICULTIES

- Swallowing oral formulations of drugs often becomes difficult for palliative care patients.
- drugs which are available in the capsule form may be more easily swallowed using the ‘leaning forward’ technique
  - this involves bending the head down rather than tipping it back when swallowing capsules
  - when leaning the head down and forward the capsule floats to the back of the throat ready to be swallowed
  - the standard way of swallowing solid oral formulations – head is tipped back – results in the capsule floating to the front of the mouth making swallowing the capsule difficult
  - this ‘leaning forward’ technique will not work for tablets as they do not float so use the standard tilting the head back approach
  - if swallowing remains an issue consider crushing tablets or opening capsules if appropriate (do not crush slow or modified release or enteric coated solid dose forms), oral liquids or other routes e.g. subcut, intranasal, sublingual, rectal
MALIGNANT ASCITES
This is a common symptom in patients with breast, colon, endometrial, ovarian, pancreatic or gastric cancers.

Assessment
- consecutive measurements of abdominal girth
- respiratory function – shortness of breath may occur
- early fullness e.g. squashed stomach
- portable ultrasound examination

Causes
- peritoneal fluid build-up in the abdomen due to a failure of the lymph system to adequately drain
- tumour in the peritoneal cavity
- low serum albumin
- excess fluid production
- venous compression or vena cava/hepatic vein thrombosis

Management
- Symptoms usually appear at > 1L of fluid in the abdomen.
- if the prognosis is short and the symptoms are not troublesome then no action may be needed
- explanation of the problem and likely outcomes may be enough to allay fears or anxieties
- if the symptoms warrant further intervention, the bowel is not distended or the ascites is not loculated, consider paracentesis
- beware of loculation – use of ultrasound is now common
- suction may be used if the fluid is viscous, e.g. of ovarian origin
- drain no more than 2L in the first hour then drain slowly for 12 to 24 hours (to a maximum of 5L in 24 hours)
- place an ostomy bag on the site once the paracentesis needle is removed to collect any residual leaking fluid
- check biochemistry frequently
- some centres advise daily measurement of girth
- a surgical opinion, for the insertion of a peritoneo-venous shunt, may help in recurrent ascites if the patient’s life expectancy is greater than 3 months
- repeated drainage may be followed by rapid reaccumulation
- drugs
  - if the patient is fit for diuretics, give spironolactone 100 mg (or more) with or without frusemide 40 mg once daily although benefit is often extremely limited
  - for gastric stasis give a prokinetic e.g. metoclopramide
  - if there is evidence of liver capsule stretch pain use a steroid e.g. dexamethasone – see co-analgesics protocol
CENTRAL NERVOUS SYSTEM

DEPRESSION
In end-of-life care it is important to distinguish between clinical depression and profound sadness.

- depression is a pervasive sense of misery
- sadness is a normal response to loss which waxes and wanes but enjoyment and future planning are retained
- most terminally ill patients do not become clinically depressed
- prevalence is about 15% (compared with 5 to 10% in the general population), most commonly in the early cancer stages
- reaching a diagnosis of depression in terminal patients is difficult as the usual physical symptoms of depression in the otherwise well such as anorexia, weight loss, sleep disturbance are often already present in patients with malignant disease whether they are depressed or not
- the psychological symptoms are more discriminative
- asking ‘Are you depressed?’ provides a bed-side assessment of mood
- suicide is rare, however, fleeting suicidal thoughts and fluctuating ‘will to live’ in cancer patients are common and not necessarily pathological
- requests for euthanasia and/or physician assisted suicide are more common although, as for suicide, this is not limited to depressed patients
- clinical depression is under-recognised and under-treated yet it is generally very responsive to treatment
- the cause of depression is unknown but imbalances in neurotransmitters, especially serotonin, in the brain may play a part

Psychological symptoms of major depression may include
- hopelessness
- anhedonia (loss of pleasure)
- morbid guilt and shame
- worthlessness and low self esteem
- request for physician assisted euthanasia
- persisting suicidal ideation
- lowered pain threshold
- decreased attention and concentration
- cognitive slowing
- impaired memory
- indecisiveness
- early morning wakening
- ruminative negative thoughts
- nihilistic and depressive delusions
- feeling of unreality
**Risk factors**
- inadequate symptom control – unrelieved pain, nausea
- poor quality of life
- lack of social support
- past and/or family history of depression
- older age
- misinformed prognosis
- drugs
  - steroids, cytotoxics, antibiotics, anti-hypertensives, neuroleptics, sedatives
- immobility
- advanced malignant disease

**Differential diagnosis**
- adjustment/grief reaction (sadness)
- ‘vital (physiological) exhaustion’
- demoralisation (a state of existential despair, meaninglessness and hopelessness but not of anhedonia and joylessness)
- delirium/sedation
- detachment (the terminal shedding of attachments)
- ‘giving up’ (affect neutral, rational, decisive)

**Management**
- mild to moderate depression
  - support, empathy, clarification of stressors or precipitators, explanation, cognitive therapy, symptomatic relief
- severe depression
  - supportive psychotherapy plus drug therapy
  - drug therapy – antidepressants are effective in 50 to 70% of cases
    > a therapeutic trial is usually appropriate
    > if in doubt, refer to a specialist psychiatrist
    > SSRI e.g. escitalopram, sertraline, fluoxetine
    > if no response in 4-6 weeks switch to mirtazapine, venlafaxine or add low dosage tricylic (e.g nortriptyline 10-20 mg – often not tolerated)
    > SNRI e.g. venlafaxine, duloxetine*
    > other – mirtazapine
  - psychostimulants e.g. methylphenidate
    not as effective as SSRIs – may help retarded/withdrawn, frail patients for a few weeks only
    > a response may be achieved from small doses (5 to 30 mg each morning) within days either alone or in combination with an SSRI – watch for additive serotonergic effects. Modafinil may be a useful alternative to methylphenidate

Drugs that are either not available or not funded in New Zealand are marked with *
DELIRIUM
Toxic confusional states, like delirium, are common in people who are dying.
• if irreversible, may be an indication of impending death
• can be most distressing for patients, family and staff

Diagnosis
• abrupt onset
• impairment of consciousness – the primary symptom which results in:
  — disorientation (to time)
  — fear and dysphoria
  — memory impairment (short term memory)
  — reduced attention span to external stimuli
  — hyperactive (frenzy) or hypoactive (retardation, torpor) but usually mixed
    hyperactive and hypoactive motor activity
  — reversal of sleep-wake cycle
  — perceptual disturbance (illusions, hallucinations)
  — disorganised thinking (paranoia, rambling)
  — dysgraphia (difficulties with writing)
• fluctuating symptoms (‘sundowner effect’)

Causes
There are often multiple organic causes but in up to 50% of cases, specific causes
are not found, despite investigations.
• infection
• organ failure (liver, kidney) and underlying medical conditions
• drugs
  — sedatives
  — anticholinergics
  — opioids
  — benzodiazepine or alcohol withdrawal
  — steroids
• metabolic disturbances
  — dehydration
  — hypercalcaemia
  — hyponatraemia
  — hypoglycaemia
• hypoxia
• anaemia (severe)
• vitamin deficiency
• cerebral metastases
• cerebral haemorrhage
• epilepsy – post-ictal
**Predisposing/precipitating/aggravating factors**

- dementia and CNS immaturity
- pain
- fatigue
- urinary retention
- constipation
- unfamiliar excessive stimuli
- change of environment

**Management**

- treat the underlying organic causes if identifiable and treatable
- treat fever, hypoxia, anaemia, dehydration, constipation, fear and anxiety and pain if possible
- ensure there is a safe and secure environment – have adequate staffing, remove potentially dangerous objects, have the mattress on the floor
- prevent sensory over-stimulation – have a single room, minimise noise and staff changes and maintain a warm and comfortable environment
- psychological interventions
  - reassurance
  - orienting aids (clock, personal belongings, presence of a supportive family)
  - cognitive strategies (clarification, reality testing, validation and repetition during lucid periods)
  - emotional support (touch, empathy)
- drugs – use if symptoms are severe (in combination with above management)
  - antipsychotics (calm or pacify rather than sedate)
    - haloperidol is the drug of choice BUT not in AIDS delirium (HIV makes the CNS more sensitive to dopamine antagonists), hepatic encephalopathy or alcohol withdrawal where benzodiazepines only should be used (see haloperidol in drug section)

**Haloperidol regimen in acute delirium:**

Oral if compliant, subcut if not

<table>
<thead>
<tr>
<th>Initial dosage</th>
<th>Mild</th>
<th>0.5 to 1.5 mg orally</th>
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<tr>
<td></td>
<td>Severe</td>
<td>1.5 to 5 mg orally</td>
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<tr>
<td></td>
<td>Very severe</td>
<td>10 mg subcut</td>
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- repeat and titrate every 30 to 40 minutes until controlled
- maintenance – 50% of daily dose required to achieve control usually 1.5 to 20 mg/day (oral)
- add anticholinergic agent e.g. benztropine 2 mg only if extrapyramidal symptoms appear
- extrapyramidal side effects are less pronounced with the parenteral route

 - levomepromazine (methotrimeprazine)
 - risperidone
 - olanzapine
 - quetiapine
— sedatives (should not be used alone in most cases of delirium as they may aggravate symptoms, particularly if inadequate doses are used, so use with an antipsychotic)
  > benzodiazepines e.g. midazolam, clonazepam
  > barbiturates e.g. phenobarbitone
  > melatonin may be useful
— anaesthetics e.g. propofol (rarely indicated)
— drug-induced delirium
  > opioid-induced – decrease dose or change opioid
  > anticholinergic-induced – e.g. physostigmine may reverse this.

Even if the aetiology is irreversible, the symptoms of delirium may be palliated. Only 10 to 20% of patients with terminal delirium should require ongoing sedation to achieve control.

DEMENTIA
Dementia is an insidious, global deterioration of intellect (problem solving), memory (short term), and personality (coarsening) without impairment of consciousness.
• a terminal disease (albeit slow) with a median survival of 4 to 5 years
• prevalence of 10% in over 65-year-olds, 20% in over 80-year-olds and for indigenous Australians the prevalence is 3 to 5 times that of non-indigenous Australians

Types
• Alzheimer’s is the most common (50% of all dementias)
  — predominant deficit is episodic memory loss
• Vascular (30% of all dementias)
  — accompanies a history of cardiovascular events (CVA)
  — islets of retained functioning
  — language is preserved
  — dysexecutive symptoms
  — gait disturbance
• Frontotemporal (FTD)
  — associated with Motor Neurone Disease
  — disinhibition, apathy and loss of empathy
  — hyperorality and compulsive ritualistic behaviours
• Lewy Body (LBD)
  — parkinsonism
  — visual hallucinations and cognitive fluctuations
  — extreme sensitivity to antipsychotics
• Reversible dementia (15%)
  — depressive pseudodementia
  — subdural and hypothyroidism
• Others
  — Huntington, alcoholic, post traumatic brain injury, paraneoplastic, postencephalitic
Assessment

- take an extensive history
- formally assess mental state using cognitive screening tools e.g. Addenbrooke’s, Montreal Cognitive Assessment

**Behavioural and Psychological Symptoms of Dementia (BPSD)**

- delirium
  - reduced brain reserves lower deliriant threshold
- depression (treat early initially with a SSRI)
- agitation/aggression (consider low dose short term antipsychotics, benzodiazepines)
  - identify precipitants (can be difficult)
  - avoid confrontation
- anxiety
  - peak in early/mid stages
- delusions (treat with antipsychotic)
  - particularly paranoid
- hallucinations – visual (more common in LBD)
- sleep/wake cycle reversal/sundowning
- loss of insight/judgement
- wandering (60% of patients)
  - pacing and lapping (exclude akathisia)
  - (dangerous) eloping i.e. getting lost, accidents
- rejection of care
  - of food, hydration (consider artificial hydration) and hygiene

**Complications**

- eating and swallowing difficulties, cachexia
- infections – pneumonia, urinary tract
  - in pneumonias the mortality is sevenfold that of a non-dementia patient
  - treat if symptomatic, antibiotics have limited efficacy
- falls – due to impulsivity, frailty, benzodiazepines
- pain – common in very elderly (50%)
  - may present behaviourally (non verbally, crying, irritability)
- adverse reactions to drugs
  - antipsychotics – sensitivity (LBD), parkinsonism, akathisia, sedation, peripheral oedema, chest infections, accelerated cognitive decline, stroke risk (3 fold that of non-dementia patients, 1.5 fold mortality), hypotension
  - benzodiazepines – sedation, falls
**Treatment**

As curative treatment does not exist ensure that end-of-life discussions/advance directives/appointment of enduring power of attorney all happen early before loss of capacity. The environment of care is important – it should be simple, safe, involve attentive and patient staff, include support and education for family and carers, person-centred, proactive, include distractions, activities, routine, memory cues and benign maternalism.

- Mild – cholinesterase inhibitors may slow early progression
- Moderate – focus on quality of life and maintenance of function
- Severe – maximise comfort, avoid aggressive, burdensome or futile treatments, avoid enteral tube nutrition, consider a secure facility, allow a natural death (AND)

**DISORDERS OF SLEEP AND WAKEFULNESS**

Sleep disturbance in people who are dying is a frequent occurrence and it requires careful assessment and management.

- sleep patterns change with age and with illness e.g. cancer
  - a reduction of depth and continuity of sleep and an increasing propensity for day-time naps occurs
  - many cancer patients have difficulty falling and staying asleep
  - cytokines are implicated in these changes

**INSOMNIA**

This is common and distressing. It undermines coping strategies through tiredness.

**Causes**

- poor symptom control of
  - anxiety, depression, pain, urinary frequency, faecal incontinence, nausea, vomiting, delirium, cough
- environmental changes
  - admission to hospital or hospice
  - disturbance by staff or family
- fear of going to sleep and never waking up
- drugs
  - stimulants e.g. methylphenidate
  - steroids (particularly if given after noon)
  - bronchodilators
  - alcohol, caffeine
- withdrawal of benzodiazepines, alcohol or tobacco

**Management**

- symptom control of above
- establish good sleep hygiene
  - regular bedtimes
  - minimise daytime napping
  - reduce evening stimulants e.g. caffeine, alcohol
— comfortable bedding
— comfortable temperature
• relaxation techniques
• drugs
  — hypnotics
  — short acting benzodiazepines e.g. temazepam
  — longer acting benzodiazepines e.g lorazepam
  — melatonin 2 to 3 mg at night
  — sedative antidepressants e.g. nortriptyline 10 to 20 mg
  — sedating antipsychotics e.g. quetiapine 25 to 50 mg at night may be considered if insomnia is resistant to above

**DROWSINESS/HYPERSONMIA**
These are common symptoms, particularly as the end-of-life approaches.

**Causes**
• organ failure e.g. renal, hepatic, cardiac, respiratory
• delirium (hypoactive)
• metabolic disturbances e.g. hyperglycaemia, hypercalcaemia
• fatigue or ‘vital exhaustion’
• infection
• raised intracranial pressure
• drugs
  — adverse effects e.g. opioids, anticholinergics, benzodiazepines, cyclizine, levomepromazine (methotrimeprazine)

**Management**
• accurate assessment
• treat/remove causes where possible
• it may be unresolvable and be a natural part of the dying process

**SLEEP PHASE (CIRCADIAN) DISORDER**
(Delayed Sleep Phase Syndrome or Sleep-Wake Reversal)
• a dysregulation of the sleep-wake cycle
  — profound initial insomnia and
  — the inability to arise at desirable hours
• particularly associated with cerebral tumours
• presents a major burden for carers

**Management**
• shifting the circadian rhythm with behavioural strategies and bright light therapy is impractical in the terminally ill
• relief care for the family and a night nurse may be necessary as this tends to be an intractable symptom
• drugs are of limited benefit
  — sedatives e.g. benzodiazepines
  — psychostimulants e.g. methylphenidate
  — sedating antipsychotics e.g. quetiapine 25 to 200 mg at night
  — pericyazine 20 to 30 mg at night
  — melatonin 0.5 to 6 mg at night

**TERMINAL RESTLESSNESS**
This may indicate physical, psychological and/or spiritual discomfort. It is often a ‘pre-death’ event.

**Causes**
- physical discomfort
  — unrelieved pain
  — distended bladder or rectum
  — physical restraint
  — insomnia
  — uncomfortable bed or environment
- delirium (see delirium section)
- psychological discomfort
  — anger
  — fear
  — guilt
  — unfinished business
- spiritual discomfort/ distress
  — helplessness
  — hopelessness
- drugs
  — akathisia induced by dopamine antagonists e.g. metoclopramide, haloperidol

**Management**
- assess and treat/remove possible causes
- explain what’s happening to the family, patient (if appropriate) or main carers
- have the family present to reassure and support
- discuss psychological discomfort e.g. anger, fear, guilt
- drugs
  — see delirium section and anxiety and fear section
  — benzodiazepines e.g. midazolam in inadequate doses can aggravate (by disinhibition) rather than relieve restlessness in some patients
  — if levomepromazine (methotrimeprazine) with a benzodiazepine are ineffective consider phenobarbitone or dexmedetomidine
PALLIATIVE SEDATION

This is considered when all other symptom-relieving measures have failed and the patient is clearly distressed.

Reasons for palliative sedation
• terminal restlessness (see terminal restlessness)
• uncontrolled delirium (see delirium)
• severe breathlessness (see dyspnoea)
• massive haemorrhage (see haemorrhage)
• neurogenic or cardiogenic pulmonary oedema
• intractable distress

How palliative sedation is achieved
• the level of sedation should be titrated to removal of distress
• drugs
  — benzodiazepines e.g. midazolam, clonazepam
  — sedating antipsychotics e.g. levomepromazine (methotrimeprazine) (subcut 12.5 to 200 mg /24 hours)
  — barbiturates e.g. phenobarbitone (subcut 600 to 1,200 mg/24 hours)
  — dexmedetomidine – experience in palliative care is limited
  — opioids
    > BUT increasing doses may not result in increased sedation (opioids tend only to be sedating in the opioid naive) and may instead induce respiratory depression or seizures

Sedation of this type may be subject to the principle of ‘double effect’ which has the dual effects of intentional relief of suffering and increased risk of hastening death.

Palliative sedation itself has not been shown to hasten death.

FEAR AND ANXIETY

FEAR
A brief, reflexive, rational and unpleasant emotional response (being afraid) caused by anticipation or awareness of danger. A present-focused, reality-based reaction initiating avoidant behaviours. Associated with physiological and psychological arousal. May be adaptive and enhance safety, or non-adaptive.
• innate fear (pain, bleeding, being alone, odours, confined spaces, novel places)
• learned fear (dying, death, being buried alive, needles, chemotherapy)

ANXIETY
Sustained and excessive uneasiness. Future-focused, irrational, grossly exaggerated response to perceived threat to the ‘self’, to one’s existence. An intrapsychic conflict. Encourages (unsuccessful) attempts to resolve threat.
• may be a normal alerting response
• may be a symptom of a medical condition (e.g. delirium, depression, hormone-secreting tumour), or a symptom of an impending medical catastrophe
may be the result of an adverse reaction to a drug e.g. bronchodilators, steroids, methylphenidate
may be a symptom of Generalised Anxiety, Panic or Depressive disorders

**Common anxieties and fears centre around:**
- being ill
- separation from loved ones, homes or jobs
- becoming dependent on others (being a ‘nuisance’ or ‘burden’)
- losing control of physical faculties
- failing to complete life goals or obligations
- uncontrolled pain or other symptoms
- abandonment
- not knowing how death will occur
- ‘death anxiety’ (the fear of non-being)
- spirituality

**Management of Fear**
- avoid threat if possible
- forwarning and preparations
- emotional first aid
- behaviour desensitisation for phobias (a syndrome of pathological fear)
- psychotropic medications of limited effectiveness

**Management of Anxiety**
- careful listening and attention to detail
- support to maintain independence and autonomy
- honest and open discussion about the future with the patient and family at a pace that they can accommodate
- support realistic hope for the future
- provide distractions to avoid boredom and excessive self-reflection
- attend to social and financial problems
- provide focussed spiritual care if appropriate
- psychotropic drugs – may be a useful adjunct
  - benzodiazepines e.g. lorazepam can be very effective in the short term (days to weeks) but this may fade and there is a risk of tolerance and dependency
  - beta-blockers e.g. propranolol may block the peripheral symptoms and thus ease the unease
  - antidepressants e.g. escitalopram, fluoxetine may be more effective longer term than benzodiazepines
RAISED INTRACRANIAL PRESSURE

Raised intracranial pressure is a life-threatening event that needs to be carefully assessed and managed to optimise quality of life and minimise symptoms.

Symptoms
- severe headache which is worse when lying down or straining
- vomiting
- convulsions
- mental – drowsiness, delirium
- diplopia
- restlessness

Causes / Risk factors
- cerebral metastases (more common with some primaries, e.g. lung, breast, melanoma than with others, e.g. prostate)
- primary brain tumour
- abscess
- cerebro-vascular event
- sagittal sinus thrombosis
- secondary hydrocephalus following surgery

Management
If raised intracranial pressure is suspected look for papilloedema and signs of cerebral irritation. Computerised tomography or MRI may be appropriate
- raise the head of the bed
- consider cranial radiotherapy or neurosurgery for malignancy if prognosis/status warrants it
- drugs
  - dexamethasone up to 16 mg per day. Avoid doses after noon as may add to insomnia. Gradually reduce dose to minimum effective. Withdraw after 7 days if ineffective (note – some anticonvulsants can reduce effectiveness – see dexamethasone page)
  - codeine (sometimes eases head pain)
  - consider anticonvulsants particularly if seizures are present
  - consider acetazolamide 250 to 500 mg once daily to bd

CONVULSIONS
Convulsions can be distressing not only for the patient but also for the family and other carers. They should be managed effectively to reduce distress and anxiety wherever possible. It is important to have a clear history of the convulsion in order to diagnose the type (grand mal, focal, absence or status epilepticus). At times a convulsion can be mistakenly diagnosed when the true cause of loss of consciousness or absence is a syncopal attack, cardiac arrhythmia, or a transient ischaemic attack.
Causes
- previously diagnosed epilepsy, brain trauma/surgery, brain tumour/mets
- drugs
  - some lower seizure threshold e.g. phenothiazines, tricyclics
  - interactions – antiepileptics have many variable and unpredictable interactions – see individual drug pages
  - withdrawal e.g. of steroids, alcohol
- metabolic disturbance, e.g. hypoxia, hyponatraemia, hypoglycaemia

Management
Prophylaxis
- drugs
  - consider dexamethasone if related to raised intracranial pressure (primary brain tumour/metastases)
  - sodium valproate initially 100 to 200 mg bd to tds increasing every 3 days to 1 to 2 g per day
  - levetiracetam 500 mg bd initially
  - carbamazepine initially 100 to 200 mg once daily to bd increasing by 100 to 200 mg every 2 weeks to 800 to 1,200 mg per day – consider therapeutic drug monitoring of plasma concentrations
  - phenytoin 200 to 300 mg nocte – consider therapeutic drug monitoring of plasma concentrations
  - if oral route is not available consider
    > clonazepam 1 to 4 mg/24 hours by subcut infusion
    > midazolam 10 to 60 mg/24 hours by subcut infusion
    > consider the use of phenobarbitone if convulsions are not effectively managed by other agents

Grand mal convulsions or status epilepticus management
- make the patient safe, explain what is happening and reassure
- drugs
  - rectal diazepam 10 to 20 mg
  - buccal midazolam 5 to 10 mg – between the cheek and gum
  - subcut boluses of clonazepam or midazolam
  - if these measures are not effective consider the use of phenobarbitone
RESPIRATORY SYSTEM

Respiratory symptoms are among the most common at the end-of-life. Dyspnoea (breathlessness), for example, can occur in more than half of patients who are dying, and the incidence increases as death approaches. In addition cough, haemoptysis, hiccups and pleural pain are present in a considerable number of people who are dying.

DYSPNOEA (BREATHELESSNESS)

Breathlessness is one of the most common and distressing symptoms for both patients and relatives as the end-of-life approaches.

- it has a reported incidence of 29 to 74% of people near the end-of-life
- the distress caused by breathlessness should not be underestimated
- a careful evaluation of the nature of the breathlessness is important
- listening to the descriptors (the language that the patient uses to describe the sensation) of the quality and quantity of breathlessness is important in choosing management
- breathlessness will only rarely be expressed in purely physical terms
- the assessment of breathlessness should use a multidimensional approach, as with the assessment of pain
- identifying the cause(s) is an essential step in effective management

Causes

- it is often multifactorial
- it is not always possible to identify one treatable cause
- impaired performance (can be broken down further into a number of separate entities)
  - airflow obstruction
- this can be related to large airways (tumour producing either extrinsic or intrinsic obstruction, laryngeal palsy, radiation stricture)
- or smaller airways (asthma, emphysema, chronic bronchitis, lymphangitis carcinomatosis)
  - decreased effective lung volume (effusions, ascites, pneumothorax, tumour, lung collapse, infection)
  - increased lung stiffness (pulmonary oedema, lymphangitis carcinomatosis, pulmonary fibrosis, mesothelioma)
  - decreased gas exchange (as above plus pulmonary emboli, thrombotic tumour, tumour effect on pulmonary circulation)
  - pain (pleurisy, chest wall infiltration, rib/vertebral fractures)
  - neuromuscular failure (paraplegia, motor neurone disease, phrenic nerve palsy, cachexia, paraneoplastic syndromes)
  - left ventricular failure (congestive heart failure)
  - ascites/pleural effusion
- increased ventilatory demand (due to anxiety, anaemia, metabolic acidosis)
Assessment

- careful assessment of each situation to identify probable causes is an essential starting point
- pay particular attention to the descriptions the patient gives of the sensation and experience of breathlessness and ask specifically “How would you describe your breathlessness today?”
- severity and meaning for each individual is important as dyspnoea may have a variable effect on quality of life at the end-of-life, varying with the cause(s) and the individual’s perception of the meaning of the symptom
- in a broad sense, dyspnoea has at least five main components, each of which must be attended to
  - sensation (what it feels like)
  - perception (how it is viewed in the context of the illness)
  - distress (does it cause suffering or grief?)
  - response (how individuals react)
  - reporting (the language used to relay these elements)

Management

- treat/remove causes where possible with treatments that are similar to those used in general medicine
  - the cancer itself together with radiation or chemotherapy
  - the complications of cancer e.g. pleural effusions, anaemia
  - concurrent non-cancer causes e.g. heart or lung disease
- non-pharmacological management
  - psychosocial support
    > address anxiety and fear by active listening and exploration of the meaning of breathlessness
    > explanation and reassurance
    > relaxation techniques
    > relearning breathing patterns and control
    > discuss coping strategies
  - positioning
  - adaptation and energy conservation which is often most effectively undertaken with the help of occupational or physio-therapists or specialist nurses
  - physiotherapy
  - drainage of effusions or ascites
  - blood transfusion may be useful if anaemia is present and it is appropriate
  - bronchial stents, brachytherapy
  - complementary therapies e.g. aromatherapy
  - music engagement, therapy and the arts
  - draughts of fresh air using fans and open windows
- at the end-of-life non-pharmacological interventions become less effective so greater reliance on drugs is common, although both may be used together
• drugs
  — opioids (usually morphine as efficacy of others have not been studied)
    > oral/parenteral – oral seems to be more effective than subcutaneous
    > doses are usually small 2.5 to 10 mg prn
  — oxygen
    > a draught of fresh air may be as effective as oxygen so only use in hypoxic patients
    > efficacy of oxygen varies between patients but if saturations are < 90% oxygen may have some benefits
  — nebulised normal saline
  — bronchodilators (nebulised/inhaled) e.g. salbutamol
    > for patients with reversible airway obstruction
  — corticosteroids e.g. dexamethasone
    > for patients with lymphangitis carcinomatosis, bronchial obstruction or radiation pneumonitis
  — benzodiazepines (short acting) e.g. midazolam
    > in anxious or fearful patients where other methods have failed
  — antibiotics e.g. amoxicillin
    > if infection is suspected may decrease secretions
  — diuretics
    > if congestive heart failure or pulmonary oedema are present
  — anticholinergics e.g. hyoscine, glycopyrrolate
    > if secretions are bothersome – see excessive (retained) secretions

COUGH

Cough is often associated with other symptoms such as dyspnoea, wheezing or chest tightness. It is a defensive mechanism – like pain – and it can have a detrimental effect on the quality of life as it interferes with communication, food and drink intake and sleep.

Causes and treatment
• acute respiratory infection
  — antibiotic (if appropriate), physiotherapy, nebulised saline
• airways disease
  — bronchodilator e.g. salbutamol, inhaled or systemic corticosteroids, physiotherapy
• malignant obstruction (tumour)
  — as above but consider nebulised local anaesthetic
• oesophageal reflux
  — prokinetic agents e.g. metoclopramide, positioning, proton pump inhibitors e.g. pantoprazole
• salivary aspiration
  — anticholinergic agent e.g. hyoscine
- cardiovascular causes
  - usual cardiac drugs
- pulmonary oedema
- drugs which can cause cough
  - angiotensin converting enzyme inhibitors e.g. captopril – change or discontinue therapy

Management
- cough with tenacious sputum i.e. a productive cough
  - may respond to steam inhalation, nebulised saline, bronchodilators or physiotherapy
- drugs (as above and below)
  - cough suppressants e.g. codeine, pholcodine, morphine
    > may be useful in dry non-productive coughs
    > titrate dose to effect
    > may not be appropriate in productive coughs as retaining the mucus may encourage infection
  - Simple linctus
    > this is a soothing syrup which may be an effective first choice
  - paroxetine (for itch of the respiratory tract)
  - nebulised local anaesthetics e.g. lignocaine (lidocaine)
    > may be useful in intractable cough
    > patients should be warned not to eat or drink for at least an hour after using the nebuliser to avoid accidental inhalation of food or drink
    > potential to cause bronchospasm so the initial dose should always be given under medical supervision
  - oxygen
    > may be useful in cough associated with emphysema
  - corticosteroids e.g. dexamethasone, prednisone
    > often used to treat cough associated with endobronchial tumours, lymphangitis or radiation pneumonitis

HICCUP
This is a respiratory reflex characterised by spasm of the diaphragm resulting in a sudden inspiration and closure of the vocal cords. Hiccup is a most distressing symptom and should be attended to with urgency. The phrenic and vagal nerve and the brain stem are involved.

Causes
- gastric distension
- diaphragmatic irritation
- phrenic or vagal nerve irritation
- uraemia
- neurological disease affecting the medulla e.g. brain stem tumour, infarction, encephalitis
- liver disease (hepatomegaly)
Management

- remove any correctable cause
  - e.g. reduction in gastric distension with a prokinetic – metoclopramide – if not obstructed
- pharyngeal stimulation with cold water
- elevation of pCO₂ using paper bag rebreathing or breath holding
- phrenic nerve block may be considered
- drugs
  - corticosteroids e.g. dexamethasone, prednisone
  - antipsychotics e.g. haloperidol, chlorpromazine, levomepromazine (methotrimeprazine)
  - muscle relaxants e.g. baclofen
  - benztropine
  - anticonvulsants may be useful if a CNS cause is present e.g. phenytoin, valproate, carbamazepine
  - gabapentin

Several of the above may have to be tried. None are consistently reliable.

EXCESSIVE (RETAINED) SECRETIONS

This phenomenon occurs when a patient is too weak to clear respiratory secretions particularly near the end-of-life.

- air passing through these secretions produces a gurgling or rattling sound (‘death rattle’) which, although not obviously distressing to the patient may be distressing for families and carers
- reassurance that the patient is not distressed is important for families

Causes

- inability to swallow or clear secretions
  - salivary or bronchial secretions
- cessation of steroids in patients with cerebral involvement can lead to neurogenic pulmonary oedema which may not respond to the management below – consider continuation of steroids in these patients

Management

- appropriate positioning to allow postural drainage
- drugs
  - anticholinergics e.g. hyoscine butylbromide, hyoscine hydrobromide, glycopyrrolate
    > can help but are often started too late in life to effect a major change as secretions already present have to evaporate first
    > hyoscine hydrobromide may cause delirium while glycopyrrolate and hyoscine butylbromide do not get into the CNS readily
- occasionally suction is needed to remove plugs of mucus but is not always successful and should be avoided if possible
HAEMOPTYSIS
The coughing up of blood from the lungs, or haemoptysis, is often a frightening symptom for both patient and family.

Causes
It is not always possible to identify the cause and it has been suggested that up to 40% of cases remain undiagnosed.
- tumour erosion – lung or oesophagus
- infection
- pulmonary embolism
- clotting disorders

Management
- treat/remove the causes if appropriate
- if minor coughing up of blood i.e. flecks or spots of blood
  - not usually helpful to give any specific treatment but patient reassurance may help
- if the bleeding is persistent or is major
  - haemostatics such as tranexamic acid may be useful (1 to 1.5 g two to four times daily)
  - consider radiotherapy which may have some benefit
- if the bleeding is massive
  - the normal ‘life saving’ interventions of bronchoscopy and intubation are inappropriate
  - reduce the patient’s awareness, fear and anxiety with subcutaneous midazolam (2.5 to 10 mg) with or without subcutaneous morphine
  - staff should stay with the patient and family until the immediate crisis is over
SKIN

ITCH (PRURITUS)

Itching can be as unpleasant and disruptive as pain and can have just as adverse an effect on quality of life.

- nerve fibres involved in the itch process are anatomically very similar to those involved in pain with opioid receptors being involved in both pathways
- cholestatic and uraemic itch in particular are mediated via opioid receptors
- the skin can be affected by many metabolic, pharmacological, dietary, environmental and psychological factors
- an accurate history of the onset and nature of itching is essential and will help to identify a cause along with examination of the skin for signs of disease
- not all itch is histamine related
- serotonin and prostaglandins may also be involved
- both central (neuropathic) and peripheral (cutaneous) itch have been identified

Causes

- hepatic/renal disease (obstructive jaundice, cholestatic and uraemic itch)
- drug allergy
- drugs e.g. opioids, vasodilators
- endocrine disease
- iron deficiency
- lymphoma
- provocative sensory influences such as rough clothing
- parasites

Management

- treat/remove causes
- attempt to break the itch/scratch cycle by short clipping nails, wearing cotton gloves, applying paste bandages
- apply surface cooling agents with emollients e.g. 0.25 to 1% menthol in aqueous cream, tepid showers, humid environment
- avoid washing with soap and use emulsifying ointment instead and Alpha-keri™ as bath oil
- light therapy may help
- drugs
  - oral anti-histamines e.g. promethazine, cetirizine
  - bile sequestrant e.g. cholestyramine 4 to 8 g per day
  - night sedation e.g. temazepam
  - H₂ antagonists (act on histamine receptors in the skin) e.g. cimetidine 400 mg twice daily
  - NSAIDs e.g. diclofenac
  - anxiolytics e.g. benzodiazepines
— steroids e.g. dexamethasone (lymphoma itch), topical hydrocortisone
— rifampicin 150 to 300 mg per day (chronic cholestasis)
— 5HT₃ antagonists e.g. ondansetron (uraemic)
— gabapentin (uraemic)
— doxepin capsules or cream
— thalidomide
— paroxetine, mirtazapine (paraneoplastic itch)

Referral to a specialist dermatologist should be considered at an early stage if no alleviation of symptoms is obtained.

SWEATING
Sweating is an unpleasant and debilitating symptom that affects not only the patient but often indirectly, the carers as well. As with many other symptoms it can indicate physical, psychological and/or environmental disturbance.

Causes
• environmental temperature changes
• emotion
  — usually confined to the axillae, palms and soles
• lymphomas, hepatic metastases and carcinoid
  — may produce drenching night sweats
• intense pain precipitating or manifesting through anxiety and fear
• infection
• drugs
  — alcohol
  — antidepressants (especially venlafaxine)
  — opioids

Management
• treat/remove causes
• drugs
  — NSAIDs e.g. diclofenac
• act via prostaglandins in the hypothalamus
  — cimetidine 400mg to 800 mg at night
• acts on histamine receptors in skin
  — steroids e.g. dexamethasone
  — paracetamol (for night sweats)
  — gabapentin
  — glycopyrrolate topically
PRESSURE INJURY CARE
Pressure injuries occur when the blood supply is shut down by pressure e.g. from a hard bed or other surface resulting in tissue death.

Causes
• pressure on one particular part of the body
  — sitting is riskier than lying as more of a person’s weight can press on a smaller area e.g. buttocks while sitting
• sliding patients against a surface can cause damage to skin (friction) or tissue (shear)
• wetness increases the risk of pressure injury damage

Assessment
• A comprehensive assessment should include:
  — clinical history
  — pressure injury risk scale
  — skin assessment
  — mobility and activity assessment
  — nutritional assessment
  — continence assessment
  — cognitive assessment
  — assessment of extrinsic risk factors

Management
• avoid causes
• assess using appropriate ‘risk factor scale’ at regular intervals i.e. daily for high risk, weekly for low risk
• use pressure relieving aids and mattresses when these are assessed as being needed
• use aids to movement where appropriate
• discuss management with patient and home carers
• use a semipermeable adhesive dressing if at risk
• where semipermeable adhesive dressing is not practical use meticulous hygiene followed by povidone iodine spray
• higher rating pressure injuries should be treated as wounds with appropriate dressing products and techniques
• rubbing over pressure injuries should be discouraged
• turn bed-fast patients every 2 to 4 hours as appropriate
• in incontinent patients protect vulnerable skin with zinc and castor oil cream and consider catheterisation
• if nutritional state is poor, get dietary advice from a dietitian
• inform primary carers of management on discharge from in-patient facility
LYMPHOEDEMA
As lymphoedema (swelling of a limb [usually] due to fluid) cannot be cured, the aim of treatment is to achieve maximal improvement and long-term control.

Causes
- damage to the lymphatic drainage system allows fluid to build up
- the protein in the initial oedema draws more fluid out of the blood
- the protein in the fluid also encourages inflammation
- infection may occur

Management
- provide analgesia if painful
- early referral to an appropriately trained professional (usually a physiotherapist) produces best results
- success requires the patient’s full cooperation, so a simple explanation of lymph flow and the cause of swelling is essential, together with instruction on daily skin care
- infections must be cleared before commencing treatment
- gentle massage of the affected area helps to shift fluid from one area to another, local practitioners in the techniques may be available
- regular measurement of both normal and affected limbs is essential to monitor progress
- in most cases containment hosiery of an appropriate size and strength should be worn all day, complemented by specific exercises and massage if possible
- if the limb is not in a suitable shape or condition to use hosiery or if the fingers are swollen, compression bandaging or taping may be necessary for approximately two weeks
- diuretics are not usually useful (except when the patient has heart failure or hypoalbuminaemia), may be detrimental and can cause dehydration

FUNGATING WOUNDS AND TUMOURS
Fungation of wounds or tumours (smelly, exuding necrotising wounds) presents an obvious manifestation of disease that can cause major distress to patient, carers and family.
- ‘fungating’ wounds are malignant in nature and combine ulceration with proliferation
- usually seen in the area of the breast or head and neck
- as healing of the wound is rare, the aim in managing these wounds is to achieve maximum patient comfort together with a reduction in the distortion of body image
- odour is often caused by anaerobic bacterial infection of compromised tissue
- the wound may bleed as blood vessels are eroded
Causes
- primary skin tumour e.g. melanoma, squamous cell carcinoma
- invasion of nearby tissue by underlying tumour e.g. breast cancer
- metastatic involvement

Management
- ensuring that the area is as clean as possible can help to reduce smell and exudate
- many preparations are recommended for odour reduction and each practitioner will have their favourite e.g. lemon oil
- as the odour is often due to anaerobic infection, metronidazole gel applied directly to the wound can be helpful
- for excessive exudate wound dressings may be used on the advice of a local expert – disposable nappies may be an option
- bismuth idoform paraffin paste (BIPP) may help in drying up the wound and reducing odour
- many fungating wounds are painful – use systemic analgesics
- morphine injection added to a gel in a clean environment and used topically may help (0.05 to 0.1% morphine [i.e. 0.5 to 1 mg/mL] in Intrasite™ gel, metronidazole gel or KY Jelly™)
- radiotherapy, chemotherapy and hormone manipulation should be considered for some tumours
- if bleeding consider pressure with adrenaline 1:1000 soaked swabs
SYSTEMIC EFFECTS OF TERMINAL DISEASES

PARANEOPlastic syndromes
The remote effects of cancer can be classified as paraneoplastic syndromes. They are thought to be rare, affecting perhaps only 1% of people with cancer. These syndromes may be identified before the diagnosis of cancer is made.

Dermatological syndromes
There are a number of skin disorders that herald the presence of underlying malignant disease. Consultation with a specialist dermatologist is advised.
- acanthosis nigricans (treatment generally ineffective)
- dermatomyositis (treatment requires removal of the cause but symptoms may be managed with corticosteroids)
  - associated with lung, breast, ovarian, pancreatic, stomach, colorectal cancers and non-Hodgkin’s lymphoma
- acquired ichthyosis (treat the underlying cause)
- paraneoplastic pemphigus (use steroids and ciclosporin)

Metabolic syndromes
- hypercalcaemia – see hypercalcaemia section
- Cushing’s syndrome (ectopic secretion of ACTH)
- SIADH – syndrome of inappropriate antidiuretic hormone secretion
  - results in hyponatraemia which is common near the end-of-life
  - symptoms appear at plasma sodium concentrations <125 mmol/L and include stupor, coma and seizures

Neurological/psychiatric syndromes
- Lambert-Eaton myasthenic syndrome (LEMS)
  - associated with small-cell lung cancer
  - manifests as muscle weakness and fatigue
  - may respond to immunosuppression, plasmapheresis and 3,4-diaminopyridine (3,4 DAP)
- sub-acute cerebellar degeneration
  - associated with ovarian and lung cancer
- polymyositis
  - associated with non-Hodgkin lymphoma, lung cancers, bladder cancers
- motor neuropathy
  - associated with lymphoma
- peripheral neuropathy
  - associated with small-cell lung cancer
- limbic encephalitis
  - changes in mood, personality
— memory impairment (recent more than remote)
— seizures

Management
All of these syndromes are usually irreversible and treatment is largely symptomatic.

VENOUS THROMBOEMBOLISM
Venous thromboembolism (VTE) includes both deep vein thrombosis (DVT) and pulmonary embolism (PE). It is a lethal disorder that is common in people with cancer and to a lesser extent in other advanced diseases.

Diagnosis/symptoms
- PE – episodic and otherwise unexplained breathlessness or confusion, tachypnoea, and pleuritic chest pain – may be difficult to interpret in the presence of other pulmonary pathology
- DVT – pain or tenderness and swelling, increased warmth, oedema and redness
- tests such as D-Dimers are generally unhelpful in advanced cancer but Doppler scans may reveal DVTs in large veins.

Causes and risk factors
- malignant disease
- recent chemotherapy or surgery
- immobility
- malignant pelvic disease
- familial (hereditary factors)
- age (over 40)
- obesity

Management
If the patient is at risk of VTE
- take into account any risk of bleeding and expected prognosis
- discuss with the patient and family whether they want to have active prophylaxis with anti-embolism stockings and low molecular weight (LMW) heparin as appropriate, balancing risks and benefits to optimise quality of life
- if the patient is in the last few days or weeks of life then thromboprophylaxis is often not appropriate, and is not routine – the best evidence in favour of thromboprophylaxis is in potentially reversible co-existing acute conditions

Treatment for VTE (DVT – includes prevention of PE and/or recurrent thrombosis)
- anticoagulation with a LMW heparin e.g. enoxaparin should be started immediately unless there is a contraindication – the preferred option because it is more effective in VTE associated with malignancy, and if dosed properly is less likely to cause bleeding
- LMW heparin followed by warfarin or dabigatran is cheaper and perhaps more convenient, but warfarin requires blood tests (INR may be very difficult to keep stable in those with advanced disease and variable nutritional intake)
- re-assess the patient regularly to confirm the management plan is appropriate to the stage of their illness and their wishes
- if warfarin is used start at the same time as LMW heparin and continue the LMW heparin for 2 days after achieving therapeutic INR
- haemorrhagic complications occur in almost 50% of people with advanced cancer (due to drug interactions or hepatic dysfunction)

**WEAKNESS/FATIGUE**

Weakness and fatigue are amongst the most common and debilitating symptoms at or near the end-of-life.
- it is often assumed that weakness is an inevitable consequence of approaching death BUT there are many factors that may exacerbate or precipitate weakness
- careful assessment may result in interventions that can improve quality of life
- there are often two main contributing factors
  - cachexia
    > a debilitating state of involuntary weight loss complicating chronic malignant, infectious and inflammatory diseases that contributes to mortality
  - asthenia
    > fatigue or lassitude
      » easily tired and a decreased capacity to maintain adequate performance
    > generalised weakness
      » anticipatory subjective sensation of difficulty in initiating a certain activity

**Causes**

**Cancer related**
- cachexia (see cachexia section)
- decreased food intake
  - nausea, vomiting, constipation, intestinal obstruction, diarrhoea, malabsorption, ‘squashed stomach syndrome’ in hepatomegaly, tumours, ascites, mouth and throat problems including infection, poor teeth, thrush, taste alteration
- metabolic problems
  - hyponatraemia, uraemia, liver failure, hypercalcaemia, anaemia from any cause
- emotional causes
  - anxiety, depression, fear, isolation, apathy, stress
- neuromuscular damage by tumour
  - to brain, spinal cord, peripheral nerves
- paraneoplastic syndromes e.g. Lambert-Eaton myasthenic syndrome, motor neuropathy
- radiotherapy and chemotherapy
- insomnia
- depression
Non-cancer related

- **drugs**
  - long-term steroids
  - some psychotropics
  - diuretics
  - antihypertensives
  - oral hypoglycaemics
  - statins

- **neurovascular problems**
  - transient ischaemic attacks, motor neurone disease, myasthenia gravis, Parkinson’s disease, peripheral neuropathies

- **metabolic diseases**
  - diabetes mellitus, Addison’s, hyper/hypothyroidism, tuberculosis, subacute bacterial endocarditis, connective tissue disorders

**Management**

- establish and, where possible, treat or remove cause
  - review the drug regimen
  - correct metabolic abnormalities

- give dietary advice/support
  - increase calorific intake if possible and appropriate

- exercise
  - exercise may be effective particularly in fatigue caused by radiotherapy
  - limited exercise programmes have been shown to be beneficial even in those close to the end-of-life

- drug therapy
  - hormones e.g. megestrol acetate, medroxyprogesterone
    > mechanism of action is unclear but dose related weight gain, improved calorie intake and improved sense of well-being have been reported
    > effect on fatigue is thought to be minimal
  - prokinetic antiemetics e.g. metoclopramide
    > decrease nausea and vomiting, increase food intake and appetite
    > no evidence of weight gain has been reported
  - steroids e.g. dexamethasone
    > weight gain and fat deposition has been documented but with no increase in lean body mass
    > benefit may be transient
  - eicosapentaenoic acid (EPA) and nutritional support in combination with anti-inflammatory agents (COX2 inhibitors) have been used
  - stimulants e.g. methylphenidate, modafinil

Although these drugs may be effective in some patients with fatigue potential benefit should be weighed against adverse effects e.g. long-term steroids causing muscle weakness.
CACHEXIA

Cachexia can be distressing for both the patient and their family and carers. It is difficult to watch a person ‘waste away’ and is often perceived as a sign of impending death.

- cachexia [derived from the Greek kakos (bad) and hexis (condition)]
- defined as a multifactorial syndrome with ongoing loss of skeletal muscle mass that cannot be fully reversed leading to progressive functional impairment
- diagnosis – weight loss greater than 5%, or 2% in individuals already showing depletion
- develops progressively through various stages – precachexia, cachexia, and refractory cachexia.
- refractory cachexia or cancer anorexia cachexia syndrome – very advanced cancer (preterminal), active catabolism low performance status (WHO score 3 or 4), and life expectancy less than 3 months
- may complicate many chronic or end-stage diseases in addition to cancer.
- not starvation, which can be reversed with nutrition.
- distinct from age-related loss of muscle mass, primary depression, malabsorption syndromes and hyperthyroidism.

Causes

The metabolic mechanism of the progressive wasting is uncertain.

- complex metabolic and catabolic processes occur with cytokines playing a major role
- tumour initiates an inflammatory response probably mediated by tumour-derived proinflammatory cytokines (interleukin-1, interleukin-6, interferon-gamma, tumour necrosis factor-alpha)
- cancer cachexia involves inflammation, hypermetabolism, neuro-hormonal changes, and the proteolytic and lipolytic factors.
- enhanced substrate cycling (fat, carbohydrate and protein) occurs which is associated with metabolic inefficiency, weight loss and a suboptimal response to nutritional support (‘anabolic blockade’)
- neural pathways controlling energy homeostasis are disturbed (particularly the hypothalamic melanocortin system), promoting catabolic activity

Assessment

Cachexia should be considered if the patient has lost ≥ 5% of their body weight and/or has a BMI < 20 kg/m² and 3 out of the following are present

- decreased muscle strength
- fatigue or reduced physical activity
- anorexia
- low fat-free mass index (low muscle mass)
- abnormal biochemistry
  - CRP > 5 mg/L
  - IL-6 > 4 pg/ml
  - Hb < 12 g/dL
  - serum albumin < 32 g/L
Treatments

• favourite foods
• un-pressured eating
• referral to a dietician
• drugs (efficacy is minimal for most)
  — dexamethasone 4 mg/day for 5 days
  — medroxyprogesterone
  — megestrol
  — EPA (up to 2 g per day)
  — cannabinoids
  — prokinetics e.g. metoclopramide
  — antidepressants e.g. mirtazapine
  — thalidomide
  — olanzapine

ANAEMIA

A significant proportion of people with advanced or chronic disease are anaemic.
• symptomatic anaemia usually presents when the haemoglobin is below 80 g/L although, if chronic, patients may adapt to this concentration

Symptoms

• fatigue
• delirium
• dyspnoea
• dizziness (postural hypotension)
• exacerbations of angina/heart failure

Causes (often multiple)

• chronic disease (normocytic)
• haemorrhage (microcytic, low iron levels)
• bone marrow failure (pancytopenic)
• malnutrition (macrocytic, folate and iron deficiencies)
• chronic renal failure (reduced erythropoietin production)

Management

• blood transfusion
  — rarely improves symptoms significantly for any length of time BUT may be considered, prior to further active treatment or a significant family event
  — it is often easier to give a transfusion rather than deal with the negotiation involved in not treating although the latter may be more appropriate
  — time, attention to detail and information for the patient and the family are all essential in the decision making and consent process
• erythropoietin
  — expensive, not readily available and response can be slow and limited
HYPERCALCAEMIA OF MALIGNANT DISEASE

The symptoms and signs of hypercalcaemia are often insidious in their onset. It can be classified as a paraneoplastic syndrome.

- should be considered in patients who have vague symptoms
- consider appropriateness of treatment BEFORE a calcium concentration
- if the patient has a serum calcium > 2.6 mmol/L consider treatment

**Symptoms**

- thirst and dehydration
- increased urinary output
- constipation
- loss of appetite
- nausea and or vomiting
- fatigue
- pain – usually back and abdominal
- confusion, depression

**Causes**

- bone metastases
- increased bone metabolism
- decreased renal clearance of calcium
- dehydration
- enhanced absorption from the gut

**Management**

- make the diagnosis
- decide about the most appropriate course of action together with the patient, family and team
- consider stopping diuretics, vitamin D and calcium
- the aim is to provide symptom relief and reduce serum calcium to an acceptable level using minimal intervention
  - mild to moderate (serum calcium 2.6 to 3 mmol/L)
    > initially oral then, if necessary, iv rehydration
    > consider steroids
  - moderate to severe (serum calcium 3 to 3.5 mmol/L)
    > initially iv or subcut rehydration
      » 2 to 3 L normal saline/24 hours
    > then iv/subcut bisphosphonate (may take 72 hours to work)
      » pamidronate 90mg iv infusion (can be given as a subcutaneous infusion)
      » zoledronic acid 4mg iv infusion can be used but is significantly more expensive
      » calcitonin may be useful when bisphosphonates begin to fail
NUTRITION IN PALLIATIVE CARE

Good nutritional advice from a dietician improves patients’ quality of life.

- ensuring food choices that are
  - appropriate to the maintenance of quality of life
  - are not detrimental to the patient i.e. aggravate nausea, or be of a difficult texture/moisture content to swallow
  - preferred foods which may entail lifting dietary restrictions and discussing with the patient’s family that food intake is no longer for the purpose of sustaining life and fuelling bodily processes
- providing an environment that allows for social interaction around meal times i.e. central dining room, playing of music during meal times
- maintaining comfort
- providing weight gain may be appropriate initially but during the terminal phase it is not an appropriate nutrition intervention goal

In some instances it may become inappropriate to hydrate or feed a patient, these cases should be discussed by a multidisciplinary team on a case by case basis.

A dietitian can provide

- complete nutrition assessments
- nutritional care plans considering an individual’s life expectancy, treatment plan and overall functional status
- assessments of nutritional factors impairing the patient’s physical and psychological well being
- patient-centred strategies such as food fortification, meal timing/ frequency and oral nutrition support
- flexible menus by liaising with catering staff to enable inpatients to enjoy their preferred foods
- an advocate role for the patient (both for and against) regarding more aggressive forms of nutrition support on a case by case basis
- clarification for the team and the patient the place of artificial nutrition when the patient is approaching the terminal phase

ORGAN FAILURE

RENAL FAILURE

The following does not apply to patients who are being dialysed. For information on drug dosing during dialysis consult a renal specialist or drug information service.

Symptoms

- oedema (from sodium and water retention)
- restless legs (may respond to clonazepam, very low dose gabapentin)
- itch (from raised urea or phosphate)
- nausea/vomiting (from increased toxins)
- fatigue (from anaemia)
**Management**
- the same as those outlined in the relevant sections e.g. nausea/vomiting
- when pain is an issue remember that
  - morphine’s metabolite is renally cleared so use fentanyl or methadone instead (or perhaps oxycodone)
  - NSAIDs increase sodium and water retention, are nephrotoxic and if urea is raised risk of GI bleed increases so avoid

**Drug dosing**
- as the kidneys fail creatinine plasma concentrations will rise
- many labs now report an estimated glomerular filtration rate (eGFR) – there is some debate as to whether this can be used to adjust the doses of renally cleared drugs
- to calculate how well the kidneys are functioning, calculate creatinine clearance in mLs/minute using the Cockcroft and Gault equation:

\[
Cr Cl (mLs/min) = \frac{(140 - age) \times ideal \text{ body weight} \ (kg) \times 0.85 \text{ if female}}{plasma \ creatinine \ (umol/L) \times 0.8}
\]

(ideal body weight = 50kg + 0.9kg for each cm above 150cm (replace 50kg with 45 kg if female))

- the creatinine clearance is important in the dosing of renally cleared drugs e.g. gabapentin or drugs whose metabolites are renally cleared e.g. morphine (see end section)
- for drugs that are almost completely renally cleared the dose regimen is a proportion of the normal dose:

\[
\text{Adjusted dose} = \frac{calculated \ creatinine \ clearance \times \ normal \ dose}{100mL/\text{min}} \times 1
\]

**HEPATIC FAILURE**
End stage liver failure is usually seen with liver metastases, liver primary and/or past alcohol abuse/hepatitis.

**Symptoms**
- raised liver enzymes
- jaundice
- ascites
- itch
- encephalopathy
- low albumin and raised INR

**Drug dosing**
- there is no single marker for liver dysfunction but albumin concentrations and INR are a measure of how well the liver can clear drugs (its metabolic capacity)
- doses of metabolised drugs (drugs that are mainly cleared from the body by the liver rather than the kidneys i.e. approx 70% of drugs) should be adjusted in severe liver failure (albumin of < 30 g/L and an INR of > 1.2) by approximately
50% especially drugs with low therapeutic index e.g. antidepressants, antipsychotics, opioids, paracetamol, anticonvulsants, NSAIDs.

Management is the same as that outlined in the relevant sections.

**CARDIAC FAILURE**

The treatment of patients with end stage cardiac failure centres around the relief of the accompanying symptoms –
- dyspnoea
- cough
- fatigue
- immobility
- oedema

Treatment of the symptoms is the same as for other causes in palliative care.

Perhaps the most difficult part of the management of these patients is when and how to discontinue the many cardiac medications prescribed (See Deprescribing section following). As yet there is no clear evidence for the order or rate of discontinuation. Negotiation with patient, family and cardiologist may produce agreement on a process for this. Once swallowing becomes a problem consideration should be given to stopping medications.

**DEPRESCRIBING IN PALLIATIVE CARE**

Deprescribing is the process of ceasing inappropriate medications safely and effectively.
- an individualised process, focusing on the patient, and taking into account their physical function, comorbidities, preferences, and lifestyle
- an ongoing process as medicines that were initially appropriately prescribed may become inappropriate over time
- often not carried out in palliative care when it perhaps should be e.g. in cancer patients who are transitioning from curative to palliative treatment or in terminally ill patients on medications with long term benefits only e.g. statins
- important because polypharmacy may lead to an underuse of essential medications and a reluctance to start new medications and an increased risk of harm due to the ‘prescribing cascade’, where more medications are prescribed to treat the side effects of others
- always consider the time required to obtain the expected benefits from medications vs expected life expectancy

**Benefits**
- improved quality of life
- reduced pill burden
- reduced potential adverse drug reactions
- improved medication adherence
**Triggers**
- older patients presenting with falls, delirium, or cognitive impairment
- development of adverse drug reactions
- worsening physiological function (cardiac/ hepatic/ renal failure)
- transition of care moments (hospital <=> home <=> palliative care unit <=> nursing home/respite)
- end-of-life

**Enablers for deprescribing**
- fear of increased adverse effects, addiction or tolerance
- inconvenience of medication taking

**Barriers to deprescribing**
- patient reluctance
- feelings of hopelessness (e.g. ‘not worth treating anymore’).
- family pressure to continue treatment, and concern from general practitioners about stopping medications first prescribed by medical specialists.

When deprescribing a medication remember that the pharmacodynamics and pharmacokinetics of other medications may be affected; use recognised tools as a starting point –
- Beers criteria
- STOPP (Screening Tool of Older Person’s Prescriptions)
- START (Screening Tool to Alert doctors to Right Treatment)
- anticholinergic risk scale

**The deprescribing process**
- take a comprehensive medication history
- ascertain indications, compliance, and potential adverse reactions
- use medication review/reconciliation services provided by pharmacists if available
- estimate life expectancy and identify any medications unlikely to provide meaningful benefit
- set goals and create a plan e.g.
  - reduced pill burden to the patient, adverse drug reactions
  - improved symptom relief and the quality of life
- emphasise that if medications are being ceased, it is not because the patient is not worth treating, but rather that the medications are causing harm or have no benefit
- relax targets of therapy e.g. levels for blood pressure, blood sugar levels, and whether blood tests should be performed
- deprescribe one or two medications at a time, not all at once
- consider a trial withdrawal to determine continuing efficacy
- provide education around what to do if symptoms return or withdrawal reactions occur
- a multidisciplinary approach should be used, with all involved and informed in the process
**Common deprescribing**

- anti-hypertensives
  - monitor blood pressure post cessation, as many patients remain normotensive
  - beta-blockers need to be weaned slowly to prevent rebound hypertension/tachycardia. NB use caution when ceasing in heart failure
- aspirin – time to benefit usually exceeds life expectancy
- diuretics – do not deprescribe if being used for symptomatic treatment or for heart failure
- statins – no evidence of benefit in shortened life expectancy or in older patients (when used for secondary prevention)
- oral hypoglycaemics (OHGs) – short term risks of continuing treatment outweigh benefits – see diabetes hyperglycaemia/hypoglycaemia page
- bisphosphonates – no evidence in shortened life expectancy, unless used for hypercalcaemia due to malignancy or for bone pain
- complementary alternative medicines (CAMs) – no evidence of benefit, unless treating a low blood plasma concentration, or to treat a symptom (zinc for taste disturbance)
- proton pump inhibitors (PPIs) – determine indication for use, as this is often not clear. Continue use if patient is on long term steroid treatment, has a history of peptic ulcer disease, active bleeding, or GORD
- cancer directed therapies – often continued in many palliative patients to improve symptoms and quality of life (e.g. preventing tumour flare at the end-of-life)
HAEMORRHAGE
Haemorrhage is distressing for all concerned and should be treated with urgency.
• in many situations the sight of blood is indicative of impending death and many patients and families experience a significant increase in anxiety – use red towels if possible
• staff are often alarmed by haemorrhage, as they often feel helpless to ‘do’ anything to prevent it
• anticipation of bleeding is sometimes possible and can be discussed with the patient and family

Management
If the patient has been taking warfarin stop it and consider reversal with fresh frozen plasma or vitamin K. If taking other anticoagulants e.g. enoxaparin or dabigatran stop them; consult a haematologist as not reversed by vitamin K.

Haemoptysis/ENT cancers
• mild
  — reassurance
• moderate
  — radiotherapy
  — bronchoscopy if appropriate
  — laser treatment if appropriate
• severe and rapid
  — subcut midazolam and/or morphine
  — have someone stay with the patient
• severe and slower
  — suction if appropriate
  — physical touch (reassures patient)
  — drugs as for severe and rapid
• other drug therapy
  — tranexamic acid 1 to 1.5 g po two to four times daily (inhibits plasminogen activation and fibrinolysis)
  — sucralfate for oral bleeding

Upper gastro-intestinal tract
• minimise causes e.g. discontinue NSAIDs
• treat gastritis and peptic ulceration
  — drug therapy (perhaps parenterally)
• proton pump inhibitor e.g. pantoprazole
• H₂ antagonist e.g. ranitidine
• radiotherapy and/or surgery may be appropriate
Lower gastro-intestinal tract
• radiotherapy and/or surgery may be appropriate
• drug therapy
  — tranexamic acid rectally
  — rectal steroids e.g. hydrocortisone rectal foam

Haematuria
• may occur with infection so check and treat if appropriate
• radiotherapy may help if tumour is present in the urinary tract
• endoscopic surgery may be appropriate
• drug therapy
  — tranexamic acid orally (as before)

Vaginal
• often due to infection so treat with antifungals and/or antibiotics
• palliative radiotherapy may help

SPINAL CORD COMPRESSION
This is a relatively uncommon problem that requires urgent and effective management.
• it is one of the true medical emergencies in palliative care
• once paralysed 95% will not walk again

Symptoms
• pain (usually before neurological symptoms)
• weakness especially of lower limbs
• sensory disturbance
• loss of sphincter control

Management
• urgent assessment
  — history and clinical findings
  — MRI examination
• referral to radiation oncology is usually most appropriate
• as soon as the diagnosis is made or suspected
  — dexamethasone 16 mg daily, for a few days then tapered down according to symptom response
  — radiation therapy should be given concurrently

Decompressive laminectomy is rarely undertaken but should be considered as an option.
MISCELLANEOUS

DIABETES, HYPERGLYCAEMIA AND HYPOGLYCAEMIA

- The pathophysiology of diabetes in the palliative care setting (and particularly in the terminal phase) may be complex as the control of blood sugar may be lost due to insulin resistance associated with illness and also because of erratic nutritional intake.
- Certain malignancies e.g. pancreatic cancer also affect the beta cells directly
- Control of blood glucose concentrations is important in palliative care as both hyperglycaemia and hypoglycaemia may cause symptoms resulting in a loss in the quality of life
  - E.g. marked hyperglycaemia may exacerbate pre-existing cachexia – in the catabolic state insulin has an anabolic effect
- Management must balance treatment tolerability (including tolerability of blood glucose monitoring if required) with treatment efficacy and symptom control

DIABETES

Type 2 diabetes (previously called non insulin dependent diabetes (NIDDM))

- Tight control of blood glucose concentrations is not necessary, although if it is easily achievable it may increase quality of life
- Relax usual dietary restrictions and adjust insulin/hypoglycaemic agent use as appropriate
- If the patient is taking metformin consider discontinuing it to avoid the adverse effects of metformin e.g. nausea, weight loss and lactic acidosis. There may be a need to add a different drug e.g. insulin
- If the patient is taking a dipeptidyl peptidase inhibitor e.g. sitagliptin this may be continued but other ‘third line’ antidiabetic agents can be discontinued e.g. pioglitazone, dapagliflozin, acarbose
- Weight loss reduces blood glucose concentrations so requirements for antidiabetic agents may reduce as weight is lost
  - Once weight loss begins or appetite decreases, halve the dose of antidiabetic agent in previously well controlled patients
  - Reduce doses further or stop as required
- On admission to a hospice oral hypoglycaemic agents i.e. sulphonylureas will not be required unless there is an infection or other serious stress in which case
  - Monitor blood glucose concentrations every two days (after the main meal if possible) and treat hyperglycaemia if symptomatic
- Symptoms of hyperglycaemia will usually appear at blood glucose concentrations of > 15 mmol/L so treatment should begin only above this concentration (in the near terminal phase, may consider treatment if blood glucose > 20-25 mmol/L)
  - Avoid hypoglycaemia during this treatment as it may be difficult to reverse without systemic therapy especially if the patient is vomiting or not eating
  - Give a fast acting insulin analogue e.g. lispro (Humalog™), aspart (NovoRapid™) or glulisine (Apidra™) insulin 2 to 4 hourly initially
(usually for 24 hours) in doses determined by monitoring – usually 5 to 10 units BUT tailor dose to both the size of the patient and food intake

— once in the range 10 to 15 mmol/L convert to an intermediate or long acting insulin e.g. isophane insulin (Protaphane™) or glargine (Lantus™) once or twice daily injections at 75% of the 24 hour short acting dose. Chart a fast acting insulin analogue e.g. lispro (Humalog™) or glulisine (Apidra™) insulin to be used for breakthrough hyperglycaemia (post-prandially if eating).

— monitor fasting blood glucose concentrations daily for several days then twice per week

— discuss management with the patient to avoid misinterpretation

**Type 1 diabetes (previously called insulin dependent diabetes (IDDM))**

— these are the minority of patients who are on insulin.

- insulin must be continued even in the terminally ill to avoid diabetic ketoacidosis. Consider capillary beta hydroxyl-butyrate monitoring if > 1.2 mmol/L ketosis is likely and should be treated if appropriate.

- tight control is not necessary
  
  — a blood glucose concentration of 10 to 15 mmol/L is a good target unless patient is symptomatic

- if the patient is well nourished and has a steady oral intake negotiate with the patient re the following
  
  — maintain the usual dose of insulin
  
  — monitor blood glucose concentrations twice a day every 3 days
  
  — when appetite decreases, increase blood glucose concentration monitoring and decrease insulin

- if patient is vomiting, is no longer eating or has a variable appetite
  
  — use a base line long acting insulin e.g. glargine (Lantus™) daily and chart a fast acting insulin analogue e.g. lispro (Humalog™) or glulisine (Apidra™) insulin to be used for breakthrough hyperglycaemia (post-prandially if eating)

  — monitor frequently

- if the patient is near to death
  
  — discuss continuation of insulin with patient and family

**HYPERGLYCAEMIA**

**Symptoms**

- at blood glucose concentrations of < 15 mmol/L
  
  — major symptoms are rare

- at blood glucose concentrations of 15 to 40 mmol/L
  
  — dehydration, dry mouth
  
  — thirst
  
  — polyuria
  
  — lethargy
  
  — blurred vision
  
  — candidiasis
  
  — skin infection
— confusion
— at blood glucose concentrations of > 40 mmol/L
— drowsiness
— obtundation
— coma

NB Some of these symptoms may be present in terminally ill patients in the absence of high blood glucose concentrations.

**Causes**

- in diabetic patients
  - lack of insulin or hypoglycaemic agent
  - loss of dietary control
  - stress, illness
  - infection
  - myocardial infarction
  - GI motility disorders and obstruction
- in non-diabetic patients
  - malignant disease
- over 1/3rd of cancer patients will develop Type 2 diabetes (NIDDM) – an effect on metabolism
- drugs (even in non-diabetic patients)
  - corticosteroids e.g. dexamethasone, prednisone
  - diuretics (at high dose) e.g. bendrofluazide, frusemide

**Management**

- in active palliative care patients
  - closely monitor blood glucose concentrations as this may help them to retain function
- in patients who are close to death
  - aim for minimal monitoring and maximal comfort
  - ‘treat the patient rather than blood glucose concentration’
  - aim for maximum quality of life by loosening control of blood glucose and encouraging eating if appropriate
- in Type 2 diabetes (non-insulin dependent) patients
  - often rehydration will partially reverse hyperglycaemia
  - BUT insulin (often only once a day) may be necessary
- in Type 1 diabetes (insulin dependent) patients
  - give insulin at least twice a day (continue with patient’s usual regimen if possible) basing the dose on body weight and predicted carbohydrate intake
  - withdrawal of insulin in these patients will lead to diabetic ketoacidosis (acidosis, shock then death), often over a period of hours or days
  - if diabetic ketoacidosis occurs treat with rehydration and iv insulin if appropriate
- drug related monitoring of blood glucose
corticosteroids e.g. dexamethasone, prednisone
• often cause hyperglycaemia
• any patient who has taken them for longer than three weeks should have intermittent blood glucose concentration monitoring
• diabetic patients taking them should have more intense blood glucose monitoring depending on the prognosis
• monitor fasting blood glucose concentrations daily for a week then three times a week for three weeks or until stable then weekly
• in terminal patients take a fasting blood glucose concentration every two days for one week and then according to clinical status

HYPOGLYCAEMIA

Symptoms – CNS
• behaviour changes, anxiety, aggression
• confusion
• fatigue
• seizures
• loss of consciousness

Symptoms – peripheral
• palpitations
• tremor
• sweating
• hunger
• paraesthesia
• pallor
• increased heart rate

Causes
• diseases
  – insulinomas (rare)
  – autoimmune disease (rare)
  – infection (sepsis)
  – carcinoid (rare)
• failure to adhere to good glucose monitoring technique
• organ failure
  – renal, hepatic, cardiac
• diet – low food intake
• drugs
  – insulin
  – hypoglycaemic agents e.g. glipizide
  – alcohol
  – quinine
  – pentamidine
Management
• treat/remove causes where possible
• give glucose (oral or iv), glucagon
• monitor blood glucose concentrations

USING STEROIDS
Steroids are often seen as cure-all/miracle drugs in palliative care. Careful consideration should be given to initiating these drugs as they have many adverse effects. Most of the use in palliative care is for unlicensed and/or non-evidence based indications e.g. spinal cord compression, nerve compression, dyspnoea (from a number of causes), SVC obstruction and inflammation following radiation therapy, pain relief, anti-cancer hormone therapy, appetite stimulation and the enhancement of well-being.

Adverse effects
• diabetes mellitus
• osteoporosis
• avascular bone necrosis
• mental disturbances
  — insomnia, paranoid psychosis, depression, euphoria
• muscle wasting (predominantly proximal myopathy)
• peptic ulceration – not as severe as NSAID induced ulceration but of concern particularly in the elderly or patients with other risk factors
• skin thinning
• immunosuppression
  — infection – candidiasis, septicaemia
  — poor wound healing
• sodium and water retention – leading to oedema
• potassium loss
• hypertension
• Cushing’s syndrome
  — moon-like face
  — striae
  — acne

Prescribing
• a trial of 5 days at 4 to 16 mg dexamethasone (dose dependent on indication) should be considered after benefit/risk has been assessed and discussed
  — dexamethasone is the preferred drug – prescribe as a single or two morning doses (before noon) to avoid sleep disturbance
• consider gastric protection with a PPI e.g. pantoprazole particularly in the elderly
• consider blood glucose monitoring (particularly if continuing)
• higher doses may be required if the patient is taking CYP enzyme inducers e.g. phenytoin and lower doses with inhibitors e.g. fluconazole
• withdraw completely if used for less than 2 weeks and < 6 mg dexamethasone. Otherwise tail off by 2 mg every 5 to 7 days until 2 mg once daily, then by 0.5 mg every 5 to 7 day
THE LAST DAYS OR HOURS

Recognising the ending of a life may seem relatively easy or obvious but in practice the ‘diagnosis of dying’ may be challenging for individuals or teams. Signs may include:

- the patient becoming increasingly weak, sleepy, disinterested in getting out of bed, seeing anyone other than close family, less interested in surroundings, confused or agitated
- symptoms becoming more apparent and physical changes suggesting the body closing down becoming more noticeable (skin colour changes, skin temperature changes, slowing of respiration or Cheyne-Stokes respiration, involuntary twitching or moaning)

Management

- planning for the death is important
- if in an institution ensure that advance care plans indicate that the person is not for resuscitation
- ensure cultural or religious wishes are known and followed
- ensure that the patient and family are aware of the progression of disease and let them know what you expect to happen
- much anxiety near the end-of-life is engendered by a fear of the unknown so provide information about those things that are known to mitigate feelings of uncertainty
- anticipate what might happen rather than wait for a crisis
- anticipatory prescribing is considered to be best practice – analgesics, antiemetics, anxiolytics and antisecretory drugs should all be considered remembering that the oral route will probably be lost so use the subcut route

Common symptoms

Pain (see pain page)

- opioids are the predominant analgesics used
- if the oral route is not feasible then consider
  - fentanyl patches – not suitable for unstable pain but may be useful as an alternative to oral analgesic
  - subcut boluses prn or continuous infusion
  - conversion from oral to subcut is 2:1 for morphine and oxycodone i.e. 10 mg oral = 5 mg subcut

Nausea/vomiting (see nausea/vomiting page)

- not usually a great problem unless there is intestinal obstruction or it has previously not been controlled

Agitation/distress/anxiety (see fear, anxiety, delirium pages)

Non-pharmacological management

- if there are fears/worries/tensions/spiritual issues consider what has helped in the past
- consider and address constipation/urinary retention/pain
Oral/buccal drugs
- lorazepam tablets 0.5 mg to 1 mg bd
- clonazepam drops (2.5 mg/mL – 0.1 mg per drop)
- midazolam sublingually or buccally (between gum and cheek)

Subcutaneous drugs
- midazolam 10 mg over 24hrs is a usual starting dose if not on benzodiazepine previously
- clonazepam boluses may be useful

Confusion (see delirium page)

Non-pharmacological management
- look for reversible causes
- aim for minimal disruption and have familiar people in the room

Oral drugs
- haloperidol drops (2 mg/mL – 0.1 mg per drop), initiate at 1 to 2 mg prn and titrate to response (much higher doses may be required – see haloperidol page)
- in frail or elderly patients an initial dose of 0.5 to 1 mg prn may be sufficient

Subcutaneous drugs
- haloperidol by continuous infusion 1 to 10 mg over 24 hours
- boluses of 1 to 2 mg may also be used

Excess secretions (see excessive (retained) secretions page)

Non-pharmacological management
- consider position change
- it may be distressing to the family/carers rather than the patient

Drugs
- glycopyrrrolate 0.6-1.2 mg subcut over 24 hours as a starting dose may help (may increase to 2.4 mg)
- hyoscine (Scopaderm™) patch may be applied behind the ear although confusion and other anticholinergic side effects may occur
- hyoscine butylbromide may be useful 20 mg subcut followed by 30 to 60 mg by continuous subcutaneous infusion over 24 hours
- secretions may become thickened and plugs may form

AFTER DEATH REVIEW
It can be helpful for teams to review what happened in order to learn from each patient and family.
- What things went well? and What lessons have been learned that can be carried to the next person and family?
- Did the patient and family resolve all unfinished business?
- Were all opportunities to say goodbye taken?
- Was death peaceful and dignified?
- Was everything possible done to care for the family and friends?
- How could care have been improved?
- How does each of the team of professional carers feel?
PALLIATIVE CHEMOTHERAPY

- palliative (i.e. non-curative) active treatments include surgery, chemotherapy and radiotherapy
- monoclonal antibody and immunotherapy drugs are being used more commonly with effect
- signal transduction inhibitors are also being used for longer (such as EGFR, BRAF, BCR-ABL, HER2 and ALK inhibitors)
- two thirds of all chemotherapy treatments given are with ‘palliative’ intent
- the aim is the palliation of symptoms but the benefit of treatment should exceed the adverse effect on quality of life
- patients of all ages who present late with chemoresponsive tumours may benefit from chemotherapy
- a few patients will gain improved survival while others may get symptom relief or time to prepare for death
- patients need to be carefully supported medically, especially if frail at the time of treatment
- although doctors may be reluctant to give chemotherapy to very ill patients, patients are often keen to try it, even if the benefits may be small

Benefits

- an often only modest survival gain of months
- chemotherapy-induced symptoms are less disruptive to quality of life than the effects of the cancer itself
- may also improve the patient and their family’s psychological well being because ‘something is being done’
- decreased tumour bulk

Adverse effects

- terminal cancer patients who receive chemotherapy during the last months of their lives are less likely to die where they wish and are more likely to undergo invasive medical procedures
- patients may express more concern about chemotherapy-induced symptoms than about the ultimate effect of the cancer
- bone marrow failure (anaemia, neutropenia, thrombocytopenia)
- unrealistic hope
- avoidance of ‘death talks’ and preparations
- nausea / vomiting
- lethargy / fatigue
- mucositis and loss of taste
- peripheral neuropathies e.g. with vincristine
- alopecia
- diarrhoea
- constipation
- stomatitis
COMPLEMENTARY AND ALTERNATIVE THERAPIES

• There is no universally agreed definition of CAM but The World Health Organisation defines it as:

‘A broad set of health care practices that are not part of a country’s own tradition and not integrated into the dominant health care system.’ Other terms sometimes used to describe these health care practices include ‘natural medicine’, ‘non-conventional medicine’ and ‘holistic medicine’.

• Complementary and Alternative Medicines (CAM) are widely used in Australasia
• a drug history should include all medicines including CAMs
• CAM can sometimes adversely impact on conventional therapies
• CAM use may be influenced by cultural beliefs and behaviours

Health professionals unfamiliar with CAM therapies that their patients are taking should seek information from a drug information pharmacist.
QUALITY OF LIFE

The primary goal of palliative care is to optimise the quality of life for patients and their families by preventing problems, delaying their onset and reducing their severity. There are many views on the nature of quality of life but one enduring view by Calman in 1984 (see further reading) is that quality of life ‘can be defined as subjective well-being reflecting differences or gaps between hopes and expectations and current experiences.’.

The aim of care near the end-of-life is to

• provide ‘appropriate’ palliative care
• provide and maintain improvement in patients’ quality of life
• achieve a ‘good death’ for the patient and family

However, health professionals and patients often have different views on what aspects of disease and treatment are important. There are many ‘expert-derived’ tools available such as:

• McGill Quality of Life questionnaire
• Schedule for the Evaluation of Individual Quality of Life (SEIQoL)
• Missoula-VITAS quality of life index – encompasses a number of domains and is user-friendly (http://www.dyingwell.org/MVQOLI.htm). It contains questions about
  — symptoms – the level of physical discomfort and distress
  — function – perceived ability to perform accustomed functions and activities of daily living and the emotional response, experienced in relation to expectations
  — interpersonal aspects – degree of investment in personal relationships and the perceived quality of one’s relations/interactions with family and friends
  — well-being – the individual’s internal condition i.e. a sense of wellness or unease, contentment or lack of contentment
  — transcendent – degree of connection with an enduring construct, and of a meaning and purpose

It has also been suggested that there are a number of developmental milestones to be reached near the end-of-life that are helpful for practitioners and patients alike to recognise including:

• a sense of completion of worldly affairs, of relationships with the community and family and friends
• a sense of meaning about our own life and life in general
• an experience of love of self and others
• an acceptance of the finality of life – of one’s existence
• a sense of a new self (personhood) beyond personal loss
• a surrender to the transcendent, to the unknown – letting go
SPIRITUALITY

Part of the ‘task of dying’ is to address spiritual concerns. Spiritual and existential concerns are important for most people at end-of-life. Spirituality should be routinely assessed, documented and addressed just as other elements of the patient’s care are. Spiritual concerns may influence other symptoms. Spiritual care needs to be patient-led and should be a normal part of history taking and care plans at end-of-life.

- there is no universally agreed definition of spirituality. It includes the existential to the religious, means different things to different people and may involve a search for: ultimate beliefs /values; a sense of meaning/purpose in life; a sense of connectedness; identity and awareness; and for some people, faith and religion. Another suggestion is that ‘spirituality is the way individuals seek and express meaning and purpose and experience their connectedness to the moment, to self, to others, to nature, to mortality and to the significant or sacred’
- spirituality is individually determined and culturally varied
- spiritual paths include nature (garden, sea, wilderness), relationships (self, family, friends, God), aesthetic pursuits (art, poetry, music), metaphysical pursuits (silence, prayer, ritual, philosophy)
- spiritual distress/pain is that caused by the threats to the extinction of the being/person and their meaning of ‘self’. It is a similar construct to demoralisation, but not to clinical depression
- there is some agreement that religion and spirituality are different but related concepts, with religion being within the broader category of spirituality although religion has become disconnected from spirituality for some

Spirituality assessment (or discernment):
The majority of seriously ill patients are likely to want their spirituality attended to, however there are a proportion who will find this intrusive. Questions that may initiate conversations are:
- ‘Are you at peace?’
- ‘What does your illness mean to you?’
- ‘Tell me about your faith?’
- ‘How is your illness challenging your relationship with your God?’
- ‘You must be wondering “Why me”?’
- ‘Do you have a belief in an afterlife?’
- ‘What gives your life meaning?’

Alternatively a spiritual wellbeing survey may be used, for example:

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Below is a list of statements that other people with your illness have said are important. Please circle or mark one number per line to indicate your response as it applies to the past 7 days.
### Dealing with spiritual distress:

- A non-judgemental approach involving presence, compassion, and empathic and contemplative listening should be used.
- The creation of space (‘a safe place to suffer’), being with and listening to (‘to be with and to bear witness’), touch and encouraging experiencing the natural and artistic worlds are useful approaches.
- Spiritual care is generally agreed to be the role of all those involved in care, with the need to involve a specialist as important as any other aspect of health care.
- More specialised interventions include retreats, group therapy, meditation and religious rituals.
- Theological beliefs and conflicts should be referred to a chaplain/pastoral care worker.
- Ethical spiritual care is critical. Proselytizing is widely understood to be unethical.

### Negative Effects of Spirituality

Not all effects associated with spirituality in the health setting are positive. The negative aspects of spirituality are mostly to do with ‘religious spirituality’. These include punishment or abandonment by God, religious pressure, guilt, stress, afterlife questions, and malign spirit visitations. The latter, and other unexplained phenomena are quite common and need to be heard compassionately. In most cases referral to a spiritual specialist is recommended.
ADVANCE CARE PLANNING (ACP) AND ADVANCE DIRECTIVES (AD)

ACP is the process of discussion and planning for future health care in the context of anticipated deterioration of health. Not everyone will choose to participate in ACP.

Health care practitioners can become familiar with the particular legal requirements in their country, state or territory by contacting relevant guardianship authorities for up-to-date information. In Australia each state and territory has different requirements. Advance care planning

- involves the patient, health care professionals and family/carers
- incorporates the patient’s beliefs, values, culture, preferences for care, current and anticipated medical status and treatment options
- needs a competent patient to participate
- should take place early in the course of a terminal illness but can happen at anytime
- may result in
  - a conversation and shared understanding between patient and health professionals
  - documentation of an ACP plan
  - the writing of an Advance Directive (see below)
  - the appointment of an enduring power of attorney/surrogate decision-maker
- is the articulation of wishes, preferences, values and goals
- respects personal autonomy and medical reality
- should be used to inform decision-making, even in acute medical emergencies
- should be regularly reviewed and updated – it is a flexible ‘living’ document
- is open to change, revision, and cancelation
- is not confined to medical issues – may include spiritual or interpersonal issues

Barriers to ACP

- it is time consuming
- there is sometimes a reluctance to discuss death and dying and the conversation may be difficult to initiate
- some patients prefer benign paternalistic medical care
- there may be an element of misinformation about the processes/rights/law
- acute/emergency interventions may not allow for consideration of the patients history
- the ‘disability paradox’ – with age and emerging health disabilities (especially cognitive) there is a tendency to moderate the assertiveness of stated care wishes

Advance directive (AD) (‘Living Will’)

- an AD is a written or oral directive/instruction about preferences for future care
- the process for completing advance directives should be raised early in the course of an illness when the patient is competent, free of undue influence and sufficiently informed
- the existence of an AD document or conversation needs to be established
- it becomes effective if the person loses capacity
it may encompass refusal of, or consent to, a particular treatment
there is no medical obligation or duty to provide treatments not offered, not effective or unavailable
clinicians are obliged to give effect to an AD but in emergencies medical indications to save life may take priority (if AD not known about)
in some states of Australia, directives are legally binding on health professionals. There may also be the provision for patients to nominate legal proxies who can make decisions on their behalf

**Competency or capacity**
- an individual’s ability to perform a particular task at a particular point in time e.g. a decision regarding their current or future health care includes competency and capacity
- all adults can be presumed to have capacity
- competency may fluctuate depending on the issues under consideration
- the patient needs to be able to understand information relevant to the decision, to reason and deliberate, to retain the information (even for only a short time), to communicate by any means
- capacity does not necessarily imply rationality
- if capacity is not possessed decisions must be taken by others in that person’s best interests and in the least restrictive manner possible

**Legally authorised proxy/surrogate decision-maker**
- refer to guidelines available for each country, state or territory as variations in the law in different areas exist
- the preferable surrogate is a close and mature relative. It is a difficult role
All decisions must be made with the patient’s best interest in mind and tend to be conservative and life-affirming.

**Testamentary Capacity**
- this is the legal and mental ability to make or alter a valid will
- the testator must have knowledge of extent and value of their property, knowledge of their natural beneficiaries, and the ability to communicate this knowledge

**GRIEF AND LOSS**
Grief is the distressing emotional response initiated by the death of a loved and attached person, or a loss. It is a normal, adjustment process. Spontaneous recovery occurs over time for the majority.
- grief begins at loss/diagnosis
- there are no specific stages of grief. Grief is never fully resolved
- modern society is death-denying and death-defying
- symptoms include sadness, anger, waves of distress, tearfulness, initial insomnia, pining, haunting reminiscences, fleeting auditory or visual pseudo-hallucinations or a sense of presence of the departed
mournning is the behavioural responses of grieving. Culture and social norms are determinants. Mourning customs serve to organise, protect and support the grief-stricken

• grief is age-influenced. Children do not develop the capacity to appreciate the permanency of death until aged 9-10. In the elderly grief may be curtailed if the death is expected
• grief therapy may be ineffective and potentially harmful, except in distressed/complicated grievers

Complicated grief
• Intense and/or protracted ( > 1-2 years)
• It is characterised by prolonged longing and yearning for the deceased, intrusive thoughts or images, anger, guilt, emotional numbness, avoidance of reminders and difficulties redefinition
• It occurs in 10-15% of bereaved people
• It is accompanied by increased psychological and physical morbidity, substance abuse and suicide
• Risk factors include sudden, unexpected, traumatic death, pre-existing dependant or ambivalent relationship, psychological/psychiatric vulnerability, disenfranchised grief (the hidden grief of those socially unable to express their response), compounded by major depression or substance abuse

Management of grief
• ‘death talk’ (anticipatory grief) and advance care planning may mitigate/moderate grief
• early identification of those at high risk for bereavement follow-up
• support, empathy, normalisation, offer pragmatic information/education
• encouraging adaptation and restructuring of a world without the lost one, acknowledgement of the emotional ‘scar’
• short term mild hypnotic medication if marked insomnia
• specific counselling e.g. Cognitive Behavioural Therapy if complicated grief, perhaps with antidepressant medication
• cathartic expression of distress is of minimal, if any, benefit
BACLOFEN

**Class:** GABA derivative musculoskeletal muscle relaxant

**Indications:** relief of musculoskeletal spasm

**Contraindications/cautions:** epilepsy, subcut injection, psychosis, schizophrenia, depression, mania, GI ulceration, cerebrovascular disease, alcoholism, diabetes (may increase blood glucose concentrations), hypertension

**Adverse reactions:**
- common nausea, sedation, somnolence;
- less common decreased cardiac output, hypotension, GI disturbance, respiratory depression, lightheadedness, personality changes, headache, insomnia, euphoria, depression, weakness, tremor, hallucinations, dry mouth, tinnitus

**Metabolism/clearance:** mainly excreted in urine unchanged (80%) so dose adjust in renal impairment

**Interactions:**
- additive drowsiness and CNS depression with other CNS depressant drugs e.g. alcohol, benzodiazepines (e.g. clonazepam), opioids
- increased muscle relaxation with tricyclic antidepressants e.g. nortriptyline

**Dosing:**
- oral: 5 to 20 mg three to four times a day (start at 5 mg three times a day)
- subcut: not available

**Syringe driver:** only intrathecal inj available – not for subcut use

**Mechanism of action:** works in the spinal cord where it stimulates GABA-receptors which inhibit the release of glutamate and aspartate (excitatory). Also has CNS depressant actions.

**Onset:** variable – hours to weeks

**Notes:** Stopping abruptly may result in a withdrawal reaction (confusion, psychosis, tachycardia, hyperthermia and rebound spasticity).
BISACODYL

Class: laxative – stimulant

Indications: constipation

Contraindications/cautions: acute abdominal pain, intestinal obstruction

Adverse reactions: *common* abdominal cramps, diarrhoea, perianal irritation (usually with suppositories); *less common* atonic colon (on prolonged use), hypokalaemia

Metabolism/clearance: mainly excreted in faeces

Interactions:
- *decreased clinical effects of antispasmodics (e.g. hyoscine butylbromide)* may occur due to stimulant effects of bisacodyl

Dosing:
- oral: 5 to 10mg at night or 5mg twice a day
- subcut: not available
- rectal: 10mg at night

Syringe driver: not available

Mechanism of action: stimulates colonic activity via nerves in the intestinal mucosa

Onset: oral: 6 to 12 hours  rectal: 20 to 60 minutes

Notes: May be useful in opioid induced constipation especially in combination with a softener.
BUPRENORPHINE*

Class: analgesic – opioid, partial mu agonist/kappa antagonist

Indications: moderate to severe pain

Contraindications/cautions: buprenorphine hypersensitivity/allergy, use with other opioids, adverse effects such as respiratory depression may not completely respond to naloxone, COPD, use with benzodiazepines

Adverse reactions: see morphine

Metabolism/clearance: metabolised by unclear pathway

Interactions:
- additive CNS depression with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), tricyclic antidepressants (e.g. amitriptyline), other opioids, alcohol

Dosing:
- sublingual combo: not used
- subcut: not used
- patch: 5 to 20mcg/hour (each patch lasts for 7 days)

Syringe driver: compatibility unknown so best to infuse on its own. Irritancy potential is unknown.

Mechanism of action: partially stimulates mu- and blocks kappa opioid receptors in the CNS and gastrointestinal tract

Peak effect: patch: 60 hours after initial application

Onset: 11 to 21 hours

Duration: patch: 7 days

Notes:
- As buprenorphine is only a partial agonist of mu receptors and an antagonist of kappa receptors it should not be used with other opioids or within 24 hours of them as it may lead to severe opioid withdrawal.
- As patches last for 7 days and peak concentrations occur at 60 hours do not use in rapidly escalating pain.
- For acute toxicity give naloxone 2 mg and repeat as required (max 10 mg) over a prolonged time but be aware that full reversal of toxicity may not occur as buprenorphine binding to opioid receptors is high.
- Do not cut patches.
- Equivalence to other opioid data are sparse but 20 micrograms /hour patch may be equivalent to 90 mg oral morphine per day.
- It is recommended that no more than two patches be applied at the same time regardless of the patch strength.
- A new patch should not be applied to the same skin site for the subsequent three to four weeks.

Drugs that are either not available or not funded in New Zealand are marked with *
CHOLESTYRAMINE

**Class:** anion exchange resin

**Indications:** hypercholesterolaemia, pruritis due to partial biliary obstruction, diarrhoea associated with ileal resection or cholerrhoeic enteropathy

**Contraindications/cautions:** complete biliary obstruction, diabetes, nephrotic syndrome, phenylketonuria, prolonged use, constipation

**Adverse reactions:** common constipation, faecal impaction, hyperchloraemic acidosis, perianal irritation, intestinal obstruction; less common nausea, bloating

**Metabolism/clearance:** combines with bile acids and is excreted in the faeces – not absorbed

**Interactions:**
- *decreased clinical effect/toxicity of some drugs* (due to decreased absorption – see below)
- *altered concentrations of some drugs that undergo enterohepatic recycling*

**Dosing:**
oral: 4 to 16 g per day

**Syringe driver:** not available

**Mechanism of action:** binds bile acids which reduces plasma bile acid concentrations

**Onset:** pruritus: 4 to 7 days

**Notes:** As absorption of other drugs will be affected take all other drugs 1 hour before or 4 to 6 hours after cholestyramine. Sachet contents must be mixed with 100 to 150 mL of fluid before administering.
CITALOPRAM

Class: Antidepressant – SSRI (Selective Serotonin Re-uptake Inhibitor)

Indications: depression, anxiety (chronic)

Contraindications/cautions: hepatic impairment, epilepsy, bleeding disorders, abrupt withdrawal

Adverse reactions: common nausea, sweating, tremor, diarrhoea (excessive serotonin), constipation, somnolence; less common dry mouth, cough, postural hypotension, tachycardia, amnesia, taste disturbance, visual disturbances, pruritus, hyponatraemia, sexual dysfunction, QT prolongation

Metabolism/clearance: metabolism unknown

Interactions:
- additive risk of serotonin syndrome (potentially fatal syndrome – symptoms include sweating, diarrhoea, confusion) with other serotonergic drugs e.g. amitriptyline, carbamazepine, fluoxetine, paroxetine, tramadol, lithium
- increased risk of bleeding (antiplatelet effect) with anticoagulants

Dosing:
oral: 10 to 40 mg once a day
subcut/rectal: not available
Syringe driver: not available

Mechanism of action: blocks the reuptake of serotonin

Onset: depression 2 to 4 weeks anxiety or pain 3 to 7 days

Peak response: 5 to 6 weeks

Notes:
- Escitalopram is available. Doses used are approximately half.
- Doses of greater than 40 mg per day have been associated with QT interval prolongation.
CLONAZEPAM

Class: anticonvulsant – benzodiazepine

Indications (NB some may be unlicensed): epilepsy, convulsions, sedation, anxiety, agitation, restless leg syndrome, neuropathic pain, dyspnoea, hiccups, myoclonic jerks

Contraindications/cautions: avoid sudden withdrawal, respiratory depression

Adverse reactions: common fatigue, drowsiness (at higher doses); less common respiratory depression, incontinence, co-ordination problems, disinhibition, increase in salivation, confusion

Metabolism/clearance: metabolised by metabolising enzyme CYP3A mainly in the liver

Interactions:
- increased clinical effect/toxicity of clonazepam (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. clarithromycin, fluconazole, grapefruit juice, itraconazole, ketoconazole
- decreased clinical effect/toxicity of clonazepam (due to decreased blood concentrations) may occur with some CYP metabolism enzyme inducers (see above) e.g. carbamazepine, phenobarbitone, phenytoin, rifampicin, St John’s wort
- additive CNS effects with other CNS depressants e.g. opioids (e.g. morphine), phenothiazines (e.g. chlorpromazine), tricyclic antidepressants (e.g. amitriptyline), alcohol may occur with concomitant clonazepam

Dosing: sedation, anxiety, agitation, restless leg syndrome, neuropathic pain, dyspnoea, hiccups, convulsions
oral: 0.5 to 8 mg a day (1 to 2 mg a day usually adequate)
subcut: 1 to 8 mg/24 hours
rectal: not available

Syringe driver: see syringe driver compatibility table

Mechanism of action: may enhance the effect of GABA, an inhibitory neurotransmitter in the CNS

Onset: oral (seizure control) 20 to 40 minutes
Half life: > 30 hours (18 to 45 hours)

Notes:
- A long acting benzodiazepine so difficult to titrate to response.
- Benzodiazepines may reduce dyspnoea by anxiolytic and sedative effects.
- Approximate equivalent oral anxiolytic/sedative doses:
  - diazepam 5 mg
  - lorazepam 0.5 to 1 mg
  - clonazepam 0.5 mg
  - temazepam 10 mg
  - midazolam 7.5 mg
  - triazolam 0.25 mg
## Pharmacological properties of benzodiazepines

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<tr>
<th>Drug</th>
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<th>Muscle relaxant</th>
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CODEINE PHOSPHATE

Class: analgesic – opioid (metabolised to morphine)

Indications (NB some may be unlicensed): step 2 in the WHO analgesic ladder, cough, diarrhoea

Contraindications/cautions: avoid use with other opioid analgesics

Adverse reactions: as for morphine – very constipating

Metabolism/clearance: metabolised by metabolising enzyme CYP2D6 mainly in the liver to an active metabolite – morphine. Minor metabolism by 3A.

Interactions:
- decreased clinical effect/toxicity of codeine (due to decreased blood concentrations of morphine – an active metabolite) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. bupropion, fluoxetine, paroxetine (not citalopram), quinine
- additive CNS effects with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), tricyclic antidepressants (e.g. amitriptyline), other opioids, alcohol may occur with concomitant codeine
- inhibition of the antidiarrhoeal effects of codeine may occur with concomitant metoclopramide/domperidone

Dosing: pain, cough and diarrhoea:
oral: 15 to 60 mg 4 to 6 hourly (Max. 240 mg in 24 hours)
subcut: not recommended – use other opioid instead
rectal: not available

Syringe driver: available as injection but not used

Mechanism of action: metabolised to morphine and other active metabolites

Peak effect: 2 to 4 hours Duration: 4 to 8 hours

Notes:
- Combination products are not recommended.
- 10% of dose is converted to morphine in “normal” metabolisers i.e. 60 mg codeine = 6 mg morphine.
- 5 to 10% of the Caucasian population may be unable to metabolise codeine to morphine.
- Combination with other opioids is illogical.
- Dihydrocodeine slow release is available although it is not often used in palliative care.
CYCLIZINE

Class: antiemetic – antihistaminic

Indications: nausea/vomiting (including motion sickness)

Contraindications/cautions: prostatic hypertrophy, narrow angle glaucoma

Adverse reactions: common drowsiness, restlessness, dry mouth, blurred vision, constipation; less common insomnia, hallucinations (more common in elderly), cardiac arrhythmias

Metabolism/clearance: metabolised in the liver mainly to norcyclizine

Interactions:
- additive CNS effects with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), tricyclic antidepressants (e.g. amitriptyline), opioids, alcohol

Dosing:
- oral: 25 to 50 mg three times a day (cyclizine hydrochloride)
- subcut: 75 to 150 mg/24 hours (cyclizine lactate) (well diluted)
- rectal: not available

Syringe driver: see syringe driver compatibility table.

Mechanism of action: acts on the histamine receptors in the vomiting centre in the CNS and has anticholinergic properties

Peak concentration: approx 2 hours

Notes:
- Although there is a theoretical interaction with prokinetic antiemetics (prokinetics stimulate the gut while cyclizine slows it down) use together is common and may be justified on the basis of central nervous system receptors antagonism.
DEXAMETHASONE

Class: corticosteroid – glucocorticoid

Indications (NB some may be unlicensed): cerebral oedema (raised intracranial pressure), allergy/anaphylaxis, replacement, shock, collagen diseases, asthma, respiratory insufficiency, leukaemia, lymphoma, rheumatic disease, psoriasis, colitis, enteritis, hypercalcaemia of malignancy, nausea/vomiting, sweating, itch, hiccup, pain, liver capsule pain, tenesmus, increased energy, weight gain

Contraindications/cautions: infections, GI bleeding

Adverse reactions: common insomnia (decrease by giving as single dose in the morning); less common sodium/fluid retention, GI ulceration, delayed wound healing, thinning of skin (on prolonged use), muscle weakness (proximal myopathy), Cushing’s syndrome, weight gain, mania, depression, delirium, hyperglycaemia, osteoporosis

Metabolism/clearance: metabolised by metabolising enzyme CYP3A (major) mainly in the liver

Interactions:
- increased clinical effect/toxicity of dexamethasone (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. aprepitant, clarithromycin, grapefruit juice, indinavir, itraconazole, ketoconazole, nelfinavir, ritonavir, telaprevir, voriconazole
- decreased clinical effect/toxicity of dexamethasone (due to decreased blood concentrations) may occur with some CYP metabolising enzyme inducers (see above) e.g. carbamazepine, phenobarbitone, phenytoin, rifampicin, St John’s wort
- decreased clinical effect/toxicity of other drugs metabolised by CYP enzymes (due to induction of their metabolism by dexamethasone) may occur e.g. aprepitant, carbamazepine, clonazepam, diazepam, domperidone, fentanyl, itraconazole, ketoconazole, methadone, midazolam, prednisone, quetiapine, triazolam
- increased risk of GI bleed/ulceration when given with NSAIDs (e.g. diclofenac)

Dosing:
oral: 4 to 32 mg in 24 hours
subcut: 4 to 16 mg/24 hours
rectal: not available

Syringe driver: see syringe drivers BUT best given as a morning bolus by subcut injection/short infusion


Onset: 8 to 24 hours

Notes:
- Anti-inflammatory effect: 3 mg dexamethasone = 20 mg prednisone = 80 mg hydrocortisone.
- On discontinuation decrease dose slowly (taper) unless the patient has been taking it for less than five days in which case dose tapering is not necessary.
- Alteration in mood is not usually seen below 6 mg dexamethasone (40 mg prednisone) per day.
• Corticosteroid-induced insomnia responds to benzodiazepines (e.g. temazepam).
• Corticosteroid induced mood disorder is usually depression and rarely mania.
• The use of steroids in palliative care is common and sometimes, particularly at high dose, consideration should be given to the appropriateness of their use.
• The use of 0.5 to 1 mg dexamethasone in a syringe driver may reduce the risk of irritation at the subcutaneous site but adverse effects can occur even at low dose.
DICLOFENAC

Class: non-steroidal anti-inflammatory drug (NSAID)

Indications (NB some may be unlicensed): pain associated with inflammation, itch, sweating

Contraindications/cautions: GI ulceration, asthma (in sensitive patients), renal, cardiac or hepatic impairment

Adverse reactions: common GI ulceration (more common if elderly, on steroids or aspirin), diarrhoea, indigestion, nausea; less common dizziness, rash, nephrotoxicity, hepatitis, oedema, hypertension, headache, tinnitus, proctitis (rectal administration)

NB inhibits platelet aggregation – may prolong bleeding time.

Metabolism/clearance: metabolised by metabolising enzyme CYP2C9 mainly in the liver

Interactions:
- increased clinical effect/toxicity of diclofenac (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. fluconazole
- decreased clinical effect/toxicity of diclofenac (due to decreased blood concentrations) may occur with some CYP metabolism enzyme inducers (see above) e.g. phenobarbitone, phenytoin, rifampicin
- increased risk of renal toxicity and hyperkalaemia with ACE inhibitors (e.g. enalapril)
- increased risk of gastro-intestinal bleed with corticosteroids (e.g. dexamethasone)
- increased clinical effect/toxicity of lithium, digoxin, methotrexate, warfarin may occur with concomitant diclofenac so monitor
- decreased clinical effects of diuretics (e.g. furosemide), antihypertensives (e.g. propranolol) may occur with concomitant diclofenac

Dosing:
oral: 50 to 150 mg per day in three divided doses for normal release and two divided doses (sometimes just one) for long acting preparations.
subcut: inj available but not for subcut injection as too irritant
rectal: as for normal release oral

Syringe driver: not recommended

Mechanism of action: inhibits prostaglandin synthesis – prostaglandins are involved in inflammation and pain

Peak effect: oral (normal release): 0.3 to 2 hours
Duration: oral (normal release): 6 to 8 hours

Notes:
- Co-analgesic often used with opioids in bone and soft tissue pain.
- NSAID of choice in palliative care.
- Patients at risk of gastro-intestinal bleeds should be prescribed gastric protection (e.g. pantoprazole) prophylactically.
DOCUSATE

**Class:** laxative – faecal softener

**Indications:** constipation

**Contraindications/cautions:** acute abdominal pain

**Adverse reactions:** *less common* abdominal cramps, atonic colon (on prolonged use), bitter taste

**Metabolism/clearance:** absorbed from the gastrointestinal tract and excreted mainly in the bile

**Interactions:**
- *decreased clinical effect of antispasmodics (e.g. hyoscine butylbromide)* may occur with concomitant docusate

**Dosing:**
- oral: 100 to 480 mg daily (with senna 1 to 2 tabs at night – Max 4 tabs)
- subcut: not available
- rectal: 1 as required

**Syringe driver:** not available

**Mechanism of action:** thought to increase intestinal secretions and facilitate their movement into faeces producing softer stools

**Onset:** oral 1 to 3 days

**Notes:**
- As docusate has some stimulant action it should be avoided in complete intestinal obstruction, as should all stimulant laxatives.
- Not laxative of choice in opioid induced constipation as a single agent but useful in combination with a stimulant (e.g. Laxsol™) although giving a softener and a stimulant as separate tablets may be more effective.
DOMPERIDONE

Class: antiemetic – prokinetic, dopamine antagonist

Indications: dyspeptic symptom complex including gastro-oesophageal reflux oesophagitis, epigastric sense of fullness, feeling of abdominal distension, upper abdominal pain, eructation, flatulence and heartburn, nausea, vomiting

Contraindications/cautions: complete intestinal obstruction

Adverse reactions: common hyperprolactinaemia, breast tenderness, QT prolongation; less common abdominal cramps, diarrhoea, dry mouth, headache, dizziness

Metabolism/clearance: metabolised by metabolising enzyme CYP3A mainly in the liver and gut.

Interactions:
- increased clinical effect/toxicity of domperidone (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. aprepitant, clarithromycin, grapefruit juice, indinavir, itraconazole, ketoconazole, nelfinavir, ritonavir, telaprevir, voriconazole
- decreased clinical effect/toxicity of domperidone (due to decreased blood concentrations) may occur with some CYP metabolising enzyme inducers (see above) e.g. carbamazepine, phenobarbitone, phenytoin, rifampicin, St John’s wort
- decreased prokinetic effect of domperidone may occur with anticholinergic drugs (e.g. amitriptyline, hyoscine)
- additive increased risk of QT interval prolongation (cardiac adverse effect which may lead to arrhythmias) with tricyclic antidepressants (e.g. amitriptyline), flecainide, erythromycin, theophylline, methotrimeprazine (levomepromazine)

Dosing:
oral: 10 mg three times a day
subcut: not available
rectal: 10 mg supp available

Syringe driver: not available

Mechanism of action: similar to metoclopramide – blocks dopamine receptors in the upper gastrointestinal tract, chemo-receptor trigger zone (CTZ) and the CNS (minimal effect on CNS therefore less likely to cause extrapyramidal side effects than metoclopramide)

Peak concentration: 30 to 110 minutes

Notes:
- Main advantage over metoclopramide is less extrapyramidal side effects but not available in injectable form.
- Useful in nausea and vomiting associated with gastric stasis.
- The United States Federal Drug Agency has warned of domperidone induced QT interval prolongation and recommend a maximum of 30 mg in 24 hours. A risk benefit assessment should be carried out when higher doses are considered along with a baseline QT interval assessment.
DULOXETINE*

Class: serotonin and noradrenaline reuptake inhibitor antidepressant

Indications (NB some may be unlicensed): depression, anxiety, neuropathic pain

Contraindications/cautions: concurrent MAOI, hepatic failure, raised ocular pressure, severe renal impairment

Adverse reactions: common insomnia, sweating, fatigue, nausea, dry mouth, diarrhoea, palpitations, blurred vision, hypotension, syncope; less common mydriasis, suicidal ideation, hepatotoxicity, hyponatraemia, serotonin syndrome, tachycardia, dizziness, tinnitus

Metabolism/clearance: Metabolised by CYP1A2 and 2D6

Interactions:

- *increased clinical effect/toxicity of duloxetine* (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. bupropion, paroxetine (not citalopram), quinine, ciprofloxacin, ketoconazole
- *decreased clinical effect/toxicity of duloxetine* (due to decreased blood concentrations) may occur with with some CYP metabolising enzyme inducers (see above) e.g. phenobarbitone, phenytoin, rifampicin, cruciferous vegetables, smoking, BBQd food
- *additive risk of serotonin syndrome* (potentially fatal syndrome – symptoms include sweating, diarrhoea, confusion) with other serotonergic drugs e.g. carbamazepine, citalopram, tricyclic antidepressants (e.g. amitriptyline), lithium, tramadol interacts with excessive alcohol to cause hepatic injury

Dosing:

- depression oral: 60 mg/day
- anxiety oral: 30 mg to 120 mg/day
- neuropathic pain oral: 60 mg to 120 mg/day
- subcut: not available
- rectal: not available

Syringe driver: not available

Mechanism of action: increases available noradrenaline and serotonin in the CNS

Drugs that are either not available or not funded in New Zealand are marked with *
ENOXAPARIN

Class: anticoagulant – low molecular weight heparin

Indications: prophylaxis of venous thromboembolic disease post-op and bedridden patients. Treatment of venous thromboembolic disease, unstable angina and myocardial infarction, Prevention of thrombus during haemodialysis, duration of more than 30 days treatment

Contraindications/cautions: heparin allergy, active bleeding, recent haemorrhagic stroke, low platelets, renal impairment (adjust dose), spinal/epidural medication, prosthetic heart valve, history of gastrointestinal ulceration/bleed

Adverse reactions: common haemorrhage, haematoma, elevated LFTs; less common allergic reactions, skin necrosis, thrombocytopenia

Metabolism/clearance: metabolised but cleared mainly by the kidneys so adjust dose in renal failure

Interactions:
• *increased effect of enoxaparin* may occur with *other drugs that decrease blood clotting* e.g. aspirin, clopidogrel, warfarin, heparin
• *increased risk of bleeding* when combined with NSAIIDs e.g. diclofenac
• *decreased effect of enoxaparin* may occur with haemostats e.g. tranexamic acid, phytomenadione (vitamin K)

Dosing:
oral: not available
subcut: treatment (of DVT etc): 1.5 mg/kg once a day or 1 mg/kg twice a day (lower in the obese and renal failure patients)
prophylaxis: 20 to 40 mg once or twice a day

Syringe driver: not available

Mechanism of action: has high anti-Xa activity

Peak anti-Xa activity: 3 to 5 hours post inj

Notes:
• As the coagulation ability of cancer patients is altered it may be that low molecular weight heparins are a better choice in these patients than oral anticoagulants.
FENTANYL

Class: analgesic – opioid

Indications: step 3 on the WHO ladder for severe pain, anaesthetic premed

Contraindications/cautions: fentanyl hypersensitivity/allergy (not nausea/hallucinations)

Adverse reactions: see morphine – less constipating (reduce dose of laxatives when converting from morphine), perhaps less sedating and less emetogenic than other opioids

Metabolism/clearance: metabolised by metabolising enzyme CYP3A mainly in the liver

Interactions:
- increased clinical effect/toxicity of fentanyl (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. aprepitant, clarithromycin, grapefruit juice, indinavir, itraconazole, ketoconazole, nelfinavir, ritonavir, telaprevir, voriconazole
- decreased clinical effect/toxicity of fentanyl (due to decreased blood concentrations) may occur with some CYP metabolism enzyme inducers (see above) e.g. carbamazepine, phenobarbital, phenytoin, rifampicin, St John’s wort
- additive CNS depression with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), tricyclic antidepressants (e.g. amitriptyline), other opioids, alcohol

Dosing:
subcut: 50 to 300 micrograms in 24 hours initially
patch: 12.5 to 300 micrograms /hour (each patch lasts for 3 days)

Syringe driver: see syringe driver compatibility table

Mechanism of action: stimulates opioid receptors in the CNS and gastrointestinal tract

Peak effect: patch: 12 to 24 hours after initial application

Duration: patch: 72 hours (plus depot effect see later)

Notes:
- Patches are unsuitable for opioid naïve patients.
- If patient is hot, or there is a heat pad near the patch, rate of absorption may increase.
- If patch comes unstuck use Micropore™ round edges to reattach.
- For acute toxicity give naloxone 2 mg and repeat as required (max 10 mg) over a prolonged time (depot in skin – see below).
- Patches leave a depot in the skin which will carry on releasing fentanyl after removal (at least 17 hours for concentrations to drop by 50%).
- Dose adjustments should usually be done every 3 days.
- Use another opioid or the fentanyl injection subcut/sublingual/intranasal for breakthrough – for fentanyl the dose may not relate to background so start at 25 micrograms fentanyl and titrate to effect.
  - Approximate conversion is morphine (po): fentanyl (subcut/patch) = 150:1 i.e. 10 mg morphine po = 66 micrograms fentanyl subcut but in chronic use this can only be used as an estimate.
Conversion Chart:

<table>
<thead>
<tr>
<th>Oral morphine (mg/24 hours)</th>
<th>fentanyl patch (mcg/hour)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;60</td>
<td>12.5</td>
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<tr>
<td>60-134</td>
<td>25</td>
</tr>
<tr>
<td>135-224</td>
<td>50</td>
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<tr>
<td>225-314</td>
<td>75</td>
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<td>315-404</td>
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<td>405-494</td>
<td>125</td>
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<tr>
<td>495-584</td>
<td>150</td>
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<td>585-674</td>
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<td>675-764</td>
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<td>945-1,034</td>
<td>275</td>
</tr>
<tr>
<td>1,035-1,124</td>
<td>300</td>
</tr>
</tbody>
</table>
FLUCONAZOLE

Class: antifungal – triazole

Indications: fungal infections – cryptococcosis, candidiasis, prophylaxis, dermatomycoses

Contraindications/cautions: renal impairment, hepatic impairment

Adverse reactions: common gastrointestinal upset, headache; less common rash (discontinue), blood disorders, arrhythmias, dizziness, convulsions, hypokalaemia

Metabolism/clearance: mainly excreted by the kidneys (fraction excreted by the kidneys unchanged = 0.8) so care in renal failure

Interactions:

- increased clinical effect/toxicity of some drugs (see below) (due to increased blood concentrations of them) may occur due to inhibition of metabolising enzymes by fluconazole e.g. diazepam, diclofenac, gliclazide, ibuprofen, indomethacin, lansoprazole, naproxen, omeprazole, pantoprazole, phenytoin, warfarin

- decreased clinical effect of amphotericin may occur with concomitant fluconazole

Dosing:
oral: vaginal candidiasis 150 mg as a single dose
cryptococcal infections/ 200 to 400 mg once a day for 7 days
systemic candidiasis
oropharyngeal candidiasis 50 to 100 mg once a day for 7 days
prophylaxis in malignancy 50 mg once a day

subcut: not usually used subcut, iv: refer to package insert

rectal: not available

Syringe driver: not applicable

Mechanism of action: inhibits fungal cell membrane formation

Notes:

- Useful in severe or recurrent fungal infections.
- May be less likely to interact with other CYP metabolised drugs (see above) than ketoconazole.
FLUOXETINE

Class: antidepressant – SSRI (Selective Serotonin Re-uptake Inhibitor)

Indications (NB some may be unlicensed): depression and associated anxiety, bulimia nervosa, obsessive-compulsive disorder, premenstrual dysphoric disorder, neuropathic pain

Contraindications/cautions: epilepsy, bleeding disorders (decreases platelet aggregation)

Adverse reactions: common nausea, sweating, tremor, diarrhoea (excessive serotonin), taste disturbance, sexual dysfunction; less common dry mouth, cough, constipation, postural hypotension, tachycardia, somnolence, amnesia, visual disturbances, pruritus, hyponatraemia

Metabolism/clearance: metabolised by metabolising enzyme CYP2D6 mainly in the liver

Interactions:
- increased clinical effect/toxicity of fluoxetine (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. bupropion, paroxetine (not citalopram), quinine
- increased clinical effect/toxicity of some drugs (due to increased blood concentrations of them) may occur with fluoxetine due to metabolising enzyme inhibition by fluoxetine e.g. amitriptyline, codeine (decreased morphine concentrations so decreased clinical efficacy of codeine), haloperidol, metoclopramide, nortriptyline, promethazime, tamoxifen (decreased endoxifen (active metabolite) concentrations so decreased clinical effects)
- additive risk of serotonin syndrome (potentially fatal syndrome – symptoms include sweating, diarrhoea, confusion) with other serotonergic drugs e.g. carbamazepine, citalopram, tricyclic antidepressants (e.g. amitriptyline), lithium, tramadol

Dosing:
oral: 20 to 80 mg in the morning
subcut: not available
rectal: not available

Syringe driver: not available

Mechanism of action: blocks the reuptake of serotonin, a neurotransmitter, in the CNS

Onset: depression/anxiety: 2 to 4 weeks  pain: 3 to 7 days

Peak response: 5 to 6 weeks

Notes:
- Fluoxetine has a half life of 48 hours but its active metabolite (norfluoxetine) has a half life of 11 days.
- Watch for serotonin syndrome if switching antidepressants as it takes four to five half lives to clear a drug from the body i.e. 44 to 55 days for fluoxetine/norfluoxetine.
- Withdrawal symptoms on stopping fluoxetine are unlikely to occur.
- Tablets are dispersible in water allowing dosing increments of < 20 mg. Capsule contents are also dispersible in water.
GABAPENTIN

Class: anticonvulsant

Indications (NB some may be unlicensed): partial seizures, including secondarily generalised tonic-clonic seizures, initially as add-on therapy in patients who have not achieved adequate control with standard antiepileptic drugs, neuropathic pain, insomnia

Contraindications/cautions: renal disease (reduce dose), absence seizures, encephalopathy

Adverse reactions: common easy bruising (purpura), increased blood pressure, dizziness, ataxia, somnolence, blurred vision; less common fatigue, headache, anxiety, GI effects, sexual dysfunction, oedema, twitching, tremor, confusion, suicidal thoughts

Metabolism/clearance: not metabolised, mainly excreted unchanged by the kidneys (fraction excreted unchanged by the kidney = 0.8) so care and adjust dose in renal dysfunction

Interactions:
- decreased clinical effect/toxicity of gabapentin with antacids e.g. Mylanta P™ due to decreased absorption of gabapentin
- additive CNS depression with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), tricyclic antidepressants (e.g. amitriptyline), opioids, alcohol

Dosing:
- oral: epilepsy 900 to 1,800 mg/day in divided doses max 2,400 mg
  neuropathic pain 900 to 3,600 mg/day in divided doses

Syringe driver: not available

Mechanism of action: may act through effects on the synthesis of GABA in the CNS
GLYCOPHYRROLATE

Class: anticholinergic – antisecretory/antispasmodic

Indications (NB some may be unlicensed): antisecretory premedication, adjunctive peptic ulceration treatment, excess/retained secretions (‘death rattle’)

Contraindications/cautions: urinary retention, cardiac disease, glaucoma

Adverse reactions: common dry mouth, tachycardia; less common urinary retention, visual problems, dizziness, constipation, drowsiness

Metabolism/clearance: excreted in the bile and unchanged by the kidneys

Interactions:
- additive anticholinergic effects (e.g. dry mouth, urinary retention) with other drugs which have anticholinergic effects e.g. cyclizine, amitriptyline, haloperidol, phenothiazines (e.g. chlorpromazine)
- decreased clinical effect (prokinetic effects) of metoclopramide / domperidone may occur with concomitant glycopyrrolate

Dosing:
oral: not available (not absorbed orally)
subcut: 200 to 600 micrograms/24 hours
rectal: not available

Syringe driver: see compatibility chart

Mechanism of action: blocks cholinergic receptors

Initial response: (im): 30 to 45 minutes
Duration: (im): 7 hours

Notes:
- May be a useful alternative to hyoscine particularly in the elderly because it is less likely to cause CNS adverse effects as it does not readily cross the blood brain barrier.
HALOPERIDOL

Class: antipsychotic – butyrophenone

Indications (NB some may be unlicensed): psychotic disorders, acute alcoholism, intractable nausea and vomiting, neuroleptanalgesia, hiccup

Contraindications/cautions: hepatic encephalopathy, epilepsy, Parkinson’s disease, DLB

Adverse reactions: common extrapyramidal symptoms (usually at 5 to 20 mg/24 hours) e.g. oculogyric crisis, dystonia, tremor, abnormal movements, restlessness - may be less with parenteral route; less common hyperprolactinaemia, dry mouth, sedation, arrhythmias, QT prolongation

Metabolism/clearance: metabolised by metabolising enzyme CYP2D6 and 3A mainly in the liver

Interactions:
- increased clinical effect/toxicity of haloperidol (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. aprepitant, bupropion, clarithromycin, fluoxetine, grapefruit juice, itraconazole, ketoconazole, paroxetine, valproate, voriconazole
- decreased clinical effect/toxicity of haloperidol (due to decreased blood concentrations) may occur with some CYP metabolism enzyme inducers (see above) e.g. carbamazepine, phenobarbitone, phencytoin, rifampicin, St John’s wort
- increased clinical effect/toxicity of some drugs (due to increased blood concentrations of them) may occur with haloperidol due to metabolising enzyme inhibition by haloperidol e.g. amitriptyline, codeine (decreased morphine concentrations so decreased clinical efficacy of codeine), metoclopramide, nortriptyline, promethazine, tamoxifen (decreased endoxifen (active metabolite) concentrations so decreased clinical effects)
- additive CNS effects with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), tricyclic antidepressants (e.g. amitriptyline), opioids, alcohol
- enhanced extrapyramidal side-effects may occur with lithium
- additive anticholinergic effects (e.g. dry mouth, urinary retention) may occur with other drugs which have anticholinergic effects e.g. cyclizine, amitriptyline, phenothiazines

Dosing:
oral: parenteral = 3 : 2 nausea/vomiting
oral: 1.5 to 3 mg once a day oral: 1.5 to 20 mg per 24 hours
subcut: 1 to 2 mg/24 hours subcut: 1 to 15 mg/24 hours
iv: 2 to 5 mg (at 1mg/minute)

Syringe driver: see syringe driver compatibility table

Mechanism of action: nausea/vomiting - blocks dopamine receptors in the chemo-receptor trigger zone thus blocking input into the vomiting centre; delirium - may rebalance the unbalanced cholinergic/dopaminergic systems seen in delirium
Peak effect: oral: 2 to 6 hours  
im/subcut: 20 minutes
Duration: up to 24 hours
Notes:

- Useful as an antiemetic where causes of nausea and vomiting are biochemical imbalance or toxins.
- Particularly useful in opioid induced nausea and vomiting. It may be given as a single oral dose at night. Doses greater than 3 mg daily add no benefit.
- Delirium: The primary pharmacological intervention for delirium is to tranquillise (to control psychotic features). Occasionally sedation (to induce sleep) is an additional requirement. (See delirium page).
HYDROMORPHONE*

Class: analgesic – opioid

Indications (NB some may be unlicensed): step 3 on the WHO ladder for severe pain, more effective in nociceptive than in neuropathic/visceral pains, severe breathlessness, cough, diarrhoea

Contraindications/cautions: hydromorphone hypersensitivity/allergy (this doesn’t mean nausea/hallucination with opioids)

Adverse reactions: common nausea/vomiting in 10 to 30% of patients (usually transient for 1 to 5 days) – give haloperidol, constipation in 90% of patients – give a stimulant & softener laxative prophylactically, dry mouth, dizziness, sedation (usually transient and on initiation or dose increase); less common respiratory depression (high doses) – pain is an antidote – give naloxone if severe, visual problems – may see things upside down/翻着，肌阵挛性抽搐 – sign of toxicity – try a different opioid, delirium in 2% of patients – give haloperidol; rare hallucinations, hyperalgesia, raised intracranial pressure, biliary/urinary tract spasm, muscle rigidity, pruritus, pulmonary oedema, physical dependence (irrelevant in dying)

Metabolism/clearance: metabolised mainly in the liver by glucuronidation to active metabolites one of which is excreted by the kidneys so watch for accumulation in renal dysfunction

Interactions:
- additive CNS effects with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), tricyclic antidepressants (e.g. amitriptyline), other opioids, MAO inhibitors

Dosing:

**pain**
- **oral:** initially 0.25 to 1 mg 4 hourly and prn
  - prescribe rescue doses of 1/5th to 1/6th of the total 24 hour dose 4 to 6 hourly
  - there is no real maximum dose but it is usually less than 200 mg/24 hours. If it is > 40 mg/24 hours consider the aetiology of the pain and the use of co-analgesia
  - review doses regularly

  subcut: oral: subcut = 2:1
  rectal: oral: rectal = 1:1
  epidural: subcut:epidural = 10:1
  intrathecal subcut:intrathecal = 100:1

**breathlessness, cough**
- **oral:** 0.5 to 1 mg 4 hourly prn

Syringe driver: see syringe driver compatibility table

Mechanism of action: stimulates mu (and other) opioid receptors in the CNS and gastrointestinal tract

Peak effect: **oral:** 1 hour

Duration: **oral:** 4 to 5 hours
Notes:

- Tolerance to effect does occur but progressive disease is also a cause of dose fade.
- Toxicity: decrease in respiratory rate, mental status and blood pressure – give naloxone (see naloxone page).
- For conversion to morphine, oxycodone, fentanyl or methadone, see relevant pages.
- Hydromorphone can affect the ability to drive. Some patients may need to be told not to drive while taking hydromorphone. Always advise patients not to drive for several days after a dose increase.

Drugs that are either not available or not funded in New Zealand are marked with *
HYOSCINE BUTYLBROMIDE

Class: antispasmodic – gastrointestinal tract

Indications (NB some may be unlicensed): GI spasm/colic, some action as anti-emetic and antisecretory, sialorrhoea, ‘death rattle’

Contraindications/cautions: megacolon, stenosis, glaucoma, tachycardia, urinary retention

Adverse reactions: common dry mouth; less common urinary retention, tachycardia, visual problems, dizziness, constipation

Metabolism/clearance: metabolised but also some excreted unchanged by the kidneys so care in renal dysfunction

Interactions:
- additive anticholinergic effects (e.g. dry mouth, urinary retention) may occur with other drugs which have anticholinergic effects e.g. cyclizine, amitriptyline, phenothiazines (e.g. chlorpromazine)
- decreased clinical effect (prokinetic effects) of metoclopramide / domperidone may occur with concomitant hyoscine butylbromide

Dosing:
- oral: 20 mg four times a day
- subcut: 40 to 100 mg/24 hours
- rectal: not available

Syringe driver: see syringe driver compatibility table

Mechanism of action: blocks the effect of acetylcholine on gastrointestinal smooth muscle causing relaxation

Onset: oral: 1 to 2 hours subcut: 5 to 10 minutes

Duration: oral: 2 hours or less

Notes:
- May be useful with steroids in intestinal obstruction.
- Doesn’t cross the blood-brain barrier so doesn’t cause drowsiness or have a central antiemetic action.
- Only 8 to 10% absorbed orally.
HYOSCINE HYDROBROMIDE

Class: anticholinergic – antisecretory

Indications (NB some may be unlicensed): premedication for sedation/amnesia, nausea/vomiting from motion sickness, ‘death rattle’

Contraindications/cautions: elderly, urinary retention, cardiac disease, glaucoma

Adverse reactions: common dry mouth, tachycardia, hypotension (especially with morphine); less common urinary retention, visual problems, dizziness, constipation, drowsiness, hallucinations (commoner in the elderly)

Interactions:
- additive anticholinergic effects (e.g. dry mouth, urinary retention) may occur with other drugs which have anticholinergic effects e.g. cyclizine, amitriptyline, phenothiazines (e.g. chlorpromazine)
- decreased clinical effect (prokinetic effects) of metoclopramide / domperidone may occur with concomitant hyoscine

Dosing:
oral: not available
subcut (as the hydrobromide): 0.4 to 2.4 mg/24 hours (usually 0.8 to 1.2 mg stat)
rectal: not available
patch: 1 patch (1.5 mg)/72 hours (behind the ear)

Syringe driver: see syringe driver compatibility table

Mechanism of action: blocks cholinergic receptors in CNS and the gastrointestinal tract

Peak response: im: 1 to 2 hours (antisecretory)
Duration: im: 8 hours

Notes:
- Thought to cross the blood brain barrier more easily then hyoscine butylbromide.
- Risk of confusion in the elderly is high.
- May be particularly useful in nausea and vomiting related to motion.
KETAMINE

Class: anaesthetic

Indications (NB some may be unlicensed): general anaesthesia (400-700 mg im), severe pain (at sub-anaesthetic doses), opioid tolerance reversal, neuropathic pain

Contraindications/cautions: hypertension, tendency to hallucinations, alcohol abuse, epilepsy

Adverse reactions: common hallucinations (see notes below), delirium, tachycardia, hypertension; less common hypotension, bradycardia, laryngospasm, diplopia, respiratory depression

Metabolism/clearance: may be metabolised in the liver by CYP metabolising enzymes. Active metabolite – norketamine

Interactions:
- additive CNS effects with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), tricyclic antidepressants (e.g. amitriptyline), opioids, alcohol

Dosing:
oral: injection has been given orally, capsules and lozenges are available as below
subcut: 100 to 500 mg in 24 hours as a ‘pulse’ over 5 days. Give a test dose of 10 mg before starting infusion.
rectal: not available

Syringe driver: see syringe driver compatibility table

Mechanism of action: in pain thought to act at NMDA receptors in the dorsal horn

Peak effect: iv: 10 to 15 minutes
Duration: iv: 15 to 30 minutes

Notes:
- May be useful in opioid tolerance/intolerance, in ‘wind-up’ (or rapidly escalating doses) and may allow a reduction in opioid dose.
- May be useful in neuropathic pain although ‘pulse’ therapy has been shown to be no better than placebo in one study.
- If hallucinations occur reduce the dose of ketamine and give a benzodiazepine (e.g. diazepam 5 mg orally, midazolam 5 mg subcutaneously) or haloperidol 2 to 5 mg orally or subcutaneously.
- Has been effective when used topically.
- ‘Pulse’ therapy (increasing subcutaneous doses over 3 to 5 days) may be sufficient to ‘reset’ the NMDA/opioid receptors. Give 100 mg/24 hours then 200 mg/24hrs then 300 mg/24hrs for 3 days then consider discontinuation.
- Oral administration usually involves lower doses e.g. 25 to 50 mg three times a day as more norketamine is produced due to first pass metabolism. Norketamine is active and may be more potent than the parent ketamine.
- Oral formulations include the injection given orally either straight or made up into a syrup (see www.palliativedrugs.com for formula), oral lozenges and oral capsules.
- Sublingual use of the injection may also be effective.
- May have a role treating severe depressive disorders.
LEVETIRACETAM

Class: anticonvulsant

Indications: seizure control

Contraindications/cautions: monitor for behavioural changes, hepatic and renal impairment

Adverse reactions: common somnolence, asthenia, infection, GI disturbance, blurred vision, hostility, pruritis

Metabolism/clearance: metabolised by hydrolysis. Fraction excreted unchanged in the urine is 0.7.

Interactions:
- increased clinical effect/toxicity of levetiracetam may occur with other drugs that are excreted by active tubular secretion e.g. probenecid
- increased clinical effect/toxicity of levetiracetam (due to increased blood concentrations) may occur with valproate
- decreased clinical effect/toxicity of levetiracetam (due to decreased blood concentrations) may occur with carbamazepine, phenobarbitone, phenytoin

Dosing:
oral: 500 mg twice daily initially (reduce in renal impairment)
sc: not available
rectal: not available

Syringe driver: not available

Mechanism of Action: inhibits Ca$^{2+}$ currents and reduces the release of Ca$^{2+}$ from intraneuronal stores. Reverses the reductions in GABA- and glycine-gated currents induced by zinc and β-carbolines.

Onset: peak concentrations at 1.5 hours
LEVOMEPROMAZINE (METHOTRIMEPRAZINE)

Class: antipsychotic/neuroleptic – phenothiazine

Indications (NB some may be unlicensed): psychosis, severe ‘terminal’ pain with anxiety/distress/restlessness, schizophrenia, with other analgesics for pain, anxiety and distress, nausea/vomiting

Contraindications/cautions: hepatic dysfunction, encephalopathy, Parkinson’s disease, DLB

Adverse reactions: common somnolence, postural hypotension, sedation; less common dry mouth, hypotension, extrapyramidal side effects (long term high dose usually)

Metabolism/clearance: metabolised by sulphonation then glucuronidation. Metabolites may be active and are excreted by the kidneys so care in renal dysfunction. May inhibit CYP2D6.

Interactions:
- increased clinical effect/toxicity of some drugs (due to increased blood concentrations of them) may occur with levomepromazine (methotrimeprazine) due to metabolising enzyme inhibition by levomepromazine (methotrimeprazine) e.g. amitriptyline, codeine (decreased morphine concentrations so decreased clinical efficacy of codeine), fluoxetine, nortriptyline, oxycodone, paroxetine, promethazine
- additive CNS effects with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), other phenothiazines (e.g. chlorpromazine), tricyclic antidepressants (e.g. amitriptyline), opioids, alcohol
- additive increased risk of QT interval prolongation (cardiac adverse effect which may lead to arrhythmias) with tricyclic antidepressants (e.g. amitriptyline), flecainide, erythromycin, theophylline, domperidone

Dosing:
- pain, restlessness, distress, delirium nausea/vomiting
- oral: 6.25 to 50 mg every 4 to 8 hours 6.25 to 12.5 mg daily
- subcut: 6.25 to 200 mg/24 hours 6.25 to 12.5 mg/24 hours
- rectal: not available

Syringe driver: dilute with 0.9% sodium chloride – see syringe driver compatibility table

Mechanism of action: suppresses sensory impulses in the CNS via various neuro-transmitters.

Onset: im/?subcut (analgesia) 20 to 40 minutes

Duration: im/?subcut 12 to 24 hours  

Half life: 15 to 30 hours

Notes:
- Only phenothiazine with analgesic properties.
- Doses of less than 25 mg/24 hours are associated with minimal sedation.
- Benztropine 2 mg may be useful in alleviating extrapyramidal side effects.
- May be a useful option in patients with multiple symptoms.
- For smaller doses disperse tablets in water and give a fraction of it.
LOPERAMIDE

Class: antidiarrhoeal – peripheral opioid receptor agonist

Indications: diarrhoea, reduce number of stools in ileostomy and colostomy patients

Contraindications/cautions: diarrhoea due to infection or antibiotics

Adverse reactions: common flatulence, constipation, abdominal distension, abdominal pain, bloating; less common giddiness, dry mouth

Metabolism/clearance: transported out of cells by P-glycoprotein which stops it crossing the blood-brain barrier. Metabolised by oxidation but 50% excreted unchanged in faeces

Interactions:
- decreased clinical effect of loperamide with prokinetics
  e.g. metoclopramide /domperidone
- CNS adverse effects may occur with P-glycoprotein inhibitors
  e.g. grapefruit juice, itraconazole, ketoconazole, tamoxifen

Dosing:
oral: 2 mg after each loose stool (max. of 16 mg/24 hours)
subcut: not available
rectal: not available

Syringe driver: not available

Mechanism of action: binds to opioid receptors in gastrointestinal tract. May also affect cholinergic receptors.

Onset: 1 to 3 hours

Notes:
- May not be of benefit if patient is already taking morphine.
- Absorbed but doesn’t normally cross the blood-brain barrier BUT may become active in the CNS as an opioid if given with P-glycoprotein inhibitors e.g. itraconazole.
LORAZEPAM

Class: anxiolytic – short acting benzodiazepine

Indications (NB some may be unlicensed): anxiety, insomnia, premedication, muscle spasm, nausea/vomiting (anxiety related)

Contraindications/cautions: respiratory failure

Adverse reactions: common sedation, dizziness, unsteadiness; less common respiratory depression (high dose), disorientation, depression, disinhibition, amnesia, excitement

Metabolism/clearance: Mainly metabolised by glucuronidation

Interactions:
- additive CNS effects with other CNS depressants e.g. other benzodiazepines (e.g. midazolam), phenothiazines (e.g. chlorpromazine), tricyclic antidepressants (e.g. amitriptyline), opioids, alcohol

Dosing:
oral: anxiety 1 to 3 mg/day in 2 to 3 doses (max. 10 mg/24 hours)
       insomnia 1 to 2 mg at bedtime

subcut: injection available (unregistered) but difficult to obtain

rectal: not available

Syringe driver: not available

Mechanism of action: may enhance the effect of GABA, an inhibitory neurotransmitter in the CNS

Onset: oral: 20 to 30 minutes sublingual: shorter onset

Duration: oral: 6 to 8 hours Half life: 10 to 20 hours

Notes:
- Lorazepam is a short acting benzodiazepine.
- Tablets may be tried sublingually.
- Not metabolised by metabolising enzymes CYP450 so less likely to interact with other drugs compared with other benzodiazepines.
- Theoretically most appropriate benzodiazepine to use in hepatic failure.
- For approximate equivalent oral anxiolytic/sedative doses see clonazepam page.
- For pharmacological properties of benzodiazepines see clonazepam page.
METHADONE

Class: analgesic – opioid

Indications (NB some may be unlicensed): step 3 in the WHO analgesic ladder, cough, opioid dependence

Contraindications/cautions: may accumulate as long half life

Adverse reactions: see morphine but less drowsiness, nausea and constipation. Has a long and variable half life so watch for signs of accumulation e.g. decreased respiratory rate or mental status (particularly in the elderly).

Metabolism/clearance: metabolised by metabolising enzyme CYP3A mainly in the liver. Demethylation is the major route of metabolism and metabolites are excreted by the kidney.

Interactions:

- increased clinical effect/toxicity of methadone (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. aprepitant, clarithromycin, grapefruit juice, indinavir, itraconazole, ketoconazole, nelfinavir, ritonavir, telaprevir, voriconazole

- decreased clinical effect/toxicity of methadone (due to decreased blood concentrations) may occur with some CYP metabolising enzyme inducers (see above) e.g. carbamazepine, phenobarbitone, phenytoin, rifampicin, St John’s wort

- additive CNS effects (including respiratory depression) with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), tricyclic antidepressants (e.g. amitriptyline), other opioids, alcohol

- additive increased risk of QT interval prolongation (cardiac adverse effect which may lead to arrhythmias) with other drugs that prolong it

Dosing: (and see notes)
oral: 2.5 to 5 mg twice daily initially
subcut: 50 to 75% of oral dose
rectal: not available in NZ

Syringe driver: see syringe driver compatibility table

Mechanism of action: stimulates opioid receptors in the CNS and gastrointestinal tract and also thought to act at the NMDA receptor

Onset: 0.5 to 1 hour initially

Duration: 6 to 8 hours initially then 22 to 48 hours on repeat dosing

Notes:

- May be useful in opioid rotation.

- Dose conversion ratios from other opioids is variable as individuals have differing methadone half lives and the ratio varies with dose (see next page).

- As affects NMDA receptors may prevent ‘wind up’ (rapidly escalating doses) on long term use and is useful in neuropathic pain.

- Renal and hepatic impairment are rarely a problem.

- Subcutaneous injection/infusion may be irritant.

- Some centres use low dose methadone alongside other opioids.
• In opioid naïve patients starting doses are usually 2.5 to 5 mg twice a day with 3 hourly prn breakthrough doses. Titrate dose weekly.

Conversion to methadone

Toombs/Ayonide method

• Convert total daily dose of morphine (or equivalent) to equivalent predicted total daily dose of methadone using the nomogram below.

• Divide the predicted total daily dose of methadone by 3 and give this dose 8 hourly e.g. total daily dose of 300 mg oral morphine (or equivalent) = total daily oral dose of methadone of 30 mg i.e. 10 mg 8 hourly.

• Breakthrough – methadone 1/10th the total daily methadone 2 hourly i.e. 10 mg 8 hourly breakthrough dose of 3 mg or continue with the original opioid for breakthrough.

• Based on ratios by Ayonide, 2000:

<table>
<thead>
<tr>
<th>mg oral morphine</th>
<th>ratio of morphine : methadone</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;100</td>
<td>3:1</td>
</tr>
<tr>
<td>101-300</td>
<td>5:1</td>
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<tr>
<td>301-600</td>
<td>10:1</td>
</tr>
<tr>
<td>601-800</td>
<td>12:1</td>
</tr>
<tr>
<td>801-1000</td>
<td>15:1</td>
</tr>
<tr>
<td>&gt;1001</td>
<td>20:1</td>
</tr>
</tbody>
</table>

Methadone Conversion Nomogram – Predicted Dose

METHYLPHENIDATE

Class: central stimulant – amphetamine related

Indications (NB some may be unlicensed): attention deficit hyperactivity disorder (possible restrictions), narcolepsy, depression, neurobehavioural symptoms in brain tumours /injuries

Contraindications/cautions: anxiety, glaucoma, agitation, hyperthyroidism, cardiac problems, hypertension, epilepsy

Adverse reactions: common nervousness, insomnia, tachycardia, urticarial; less common blurred vision, hallucinations, blood disorders, psychosis (very high doses), arrhythmias

Metabolism/clearance: metabolised by hydrolysis. Inactive metabolite is excreted by the kidneys.

Interactions:
- increased analgesia and decreased sedation may occur with some opioids
- hypertensive crisis may occur with concomitant MAOIs (e.g. tranylcypromine)
- decreased hypotensive effect of adrenergic blockers (e.g. terazosin) may occur with concomitant methylphenidate
- hypertension with tricyclic antidepressants (e.g. amitriptyline) may occur

Dosing: depression (max. adult dose of 1 mg/kg/24 hours)
- oral: normal release 10 to 30 mg a day (morning and mid-day)
- subcut: not available
- rectal: not available

Syringe driver: not available

Mechanism of action: acts as a stimulant in the CNS

Onset: depression 2 to 5 days

Notes:
- Patients may respond to short courses of 2 to 3 weeks then withdraw.
- Methylphenidate is occasionally used to treat opioid-induced drowsiness.
METOCLOPRAMIDE

Class: antiemetic – prokinetic

Indications: nausea and/or vomiting, restoration of tone in upper GI tract, hiccups

Contraindications/cautions: complete intestinal obstruction. Young persons (< 20 years old) are more prone to extrapyramidal side effects so use lower doses

Adverse reactions: less common tardive dyskinesia – usually on prolonged use, extrapyramidal reactions e.g. Parkinsonism, akathisia (usually at doses > 30 mg/24 hours – switch to domperidone which enters the CNS to a lesser extent), diarrhoea, restlessness

Metabolism/clearance: metabolised in the liver partially by the metabolising enzyme CYP2D6 to inactive metabolites which are mainly excreted with some parent drug by the kidneys

Interactions:

- increased clinical effect/toxicity of metoclopramide (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. bupropion, fluoxetine, paroxetine, quinine
- faster onset of action of SR morphine may occur with concomitant metoclopramide
- prokinetic activity of metoclopramide may be affected by concomitant opioids, anticholinergics e.g. hyoscine
- increased risk of extrapyramidal effects and neurotoxicity with lithium

Dosing:
oral: 10 mg three times a day (max. 0.5 mg/kg)
subcut: 30 to 60 mg over 24 hours (watch for extrapyramidal effects at > 30 mg/24 hours)
rectal: 10 mg up to three times a day

Syringe driver: see syringe driver compatibility table

Mechanism of action: blocks dopamine receptors and perhaps affects 5HT receptors in the gastro-intestinal tract (increasing peristalsis), CNS and chemoreceptor-trigger zone (CTZ)

Peak effect: oral/rectal 1 to 3 hours

Notes:

- ‘High dose’ metoclopramide may work via 5HT3 antagonism (like ondansetron) but is associated with severe extrapyramidal effects.
- Most effective for nausea/vomiting due to gastric stasis. Some clinicians believe that metoclopramide is no better than placebo as an antiemetic but is useful as a prokinetic.
- Benztropine 2 mg may be used as an antidote.
- The European Medicines Agency’s Committee recommends that metoclopramide should only be prescribed for short-term use (up to five days) and that it should only be used as a second-line.
METRONIDAZOLE

Class: antibiotic – anti-anaerobe

Indications (NB some may be unlicensed): bacterial infections, useful in controlling malodorous wounds

Adverse reactions: common GI upset, urticaria, metallic taste, furry tongue; less common drowsiness, headache, dizziness, urine darkening, blood disorders, muscle/joint pain

Metabolism/clearance: metabolised in the liver to some active and some inactive metabolites which are excreted with some parent drug by the kidneys

Interactions:
- disulfiram-like reaction (nausea, vomiting, sweating) may occur with concomitant alcohol
- increased toxicity of lithium may occur with metronidazole

Dosing:
oral: 800 mg stat then 400 mg three times a day
subcut: injection available but not usually used subcut
iv: 500 mg three times a day (infusion)
rectal: 1 g three times a day for 3 days then twice a day
topical: apply twice a day

Syringe driver: not applicable

Mechanism of action: in malodorous wounds kills anaerobes responsible for the smell.
MICONAZOLE

Class: antifungal – imidazole

Indications: fungal infection – topical, oral, GI, vaginal

Contraindications/cautions: hepatic impairment

Adverse reactions: common oral gel – GI upset; less common oral gel – hepatitis, topical/vaginal-burning, itching

Metabolism/clearance: metabolised by the liver

Interactions: Oral gel/vaginal preparations (absorption is likely)
• decreased clinical effect of amphotericin may occur with miconazole
• may affect INR of patients taking warfarin. Monitor even if only using oral gel

Dosing:
mouth (topical): 50 mg four times a day for 7 days
subcut: not available
rectal: not available
topical: apply twice a day
vaginal: use at night

Syringe driver: not available

Mechanism of action: increases fungal cell membrane permeability
MICROLAX™/MICOLETTE™
(Sodium citrate 450mg, sodium lauryl sulphoacetate 45mg, sorbitol 3.125g, sorbic acid 5mg, water to 5mL)

Class: rectal laxative – stimulant, faecal softener and osmotic

Indications: constipation, bowel evacuation

Dosing:
oral: not available
subcut: not available
rectal: 1 tube as required

Syringe driver: not available

Mechanism of Action: may stimulate colonic activity via nerves in the intestinal mucosa (sodium citrate) and increased fluid uptake by stools thus softening them (sodium lauryl sulphoacetate, sorbitol)

Onset: almost immediate
MIDAZOLAM

Class: sedative – benzodiazepine

Indications (NB some may be unlicensed): sedation, anaesthetic induction agent, hiccups, epilepsy, muscle spasm, dyspnoea, insomnia

Contraindications/cautions: avoid sudden withdrawal, respiratory depression

Adverse reactions: common fatigue, drowsiness, amnesia; less common respiratory depression (high dose), aggression, confusion, hypotension

Metabolism/clearance: metabolised by metabolising enzyme CYP3A (major) mainly in the liver

Interactions:

- increased clinical effect/toxicity of midazolam (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. clarithromycin, fluconazole, grapefruit juice, itraconazole, ketoconazole
- decreased clinical effect/toxicity of midazolam (due to decreased blood concentrations) may occur with some CYP metabolism enzyme inducers (see above) e.g. carbamazepine, phenobarbitone, phenytoin, rifampicin, St John’s wort
- additive CNS effects with other CNS depressants e.g. other benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), tricyclic antidepressants (e.g. amitriptyline), opioids, alcohol

Dosing:
oral: 7.5 to 15 mg at bed-time
subcut: 5 to 60 mg/24 hours (up to 150 mg in sedation at the end-of-life)
rectal: not available

Syringe driver: see syringe driver compatibility table

Mechanism of action: may enhance the effect of GABA, an inhibitory neurotransmitter in the CNS

Peak concentrations: oral 20 to 50 min, subcut 5 to 10 min, iv 2 to 3 mins

Duration: 15 minutes to several hours, Half life: 2 to 5 hours

Notes:
- Midazolam is a very short acting benzodiazepine so dose titration to response is easier than with longer acting benzodiazepines e.g. clonazepam.
- iv administration can result in hypotension and transient apnoea.
- Benzodiazepines may reduce dyspnoea by anxiolytic and sedative effects.
- For approximate equivalent oral anxiolytic/sedative doses see clonazepam page.
- For pharmacological properties of benzodiazepines and other hypnotics see clonazepam page.
- May be used buccally.
MIRTAZAPINE

**Class:** antidepressant – central presynaptic alpha 2 and 5HT antagonist

**Indications (NB some may be unlicensed):** major depression, nausea

**Contraindications/cautions:** bipolar depression, epilepsy, cardiac disease, prostatic hypertrophy, diabetes, abrupt withdrawal

**Adverse reactions:** *common* increased appetite, dizziness, headache, dry mouth; *less common* convulsions, tremor, nightmares, mania, syncope, hyponatraemia, nausea

**Metabolism/clearance:** metabolised by metabolising enzyme CYP2D6, 1A2 and 3A mainly in the liver to at least one active metabolite (by CYP3A)

**Interactions:**
- *increased clinical effect/toxicity of mirtazapine* (due to increased blood concentrations of parent) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. bupropion, aprepitant, ciprofloxacin, clarithromycin, fluconazole, fluoxetine, grapefruit juice, itraconazole, ketoconazole, paroxetine, quinine
- *decreased clinical effect/toxicity of mirtazapine* (due to decreased blood concentrations of parent) may occur with some CYP metabolism enzyme inducers (see above) e.g. broccoli, carbamazepine, dexamethasone, phenobarbitone, phenytoin, prednisone, rifampicin, smoking, St John’s wort
- *additive risk of serotonin syndrome* (potentially fatal syndrome – symptoms include sweating, diarrhoea, confusion) with other serotonergic drugs e.g. amitriptyline, carbamazepine, fluoxetine, paroxetine, tramadol, lithium

**Dosing:**
- oral: 15 to 45 mg at bed-time
- subcut: not available

**Syringe driver:** not available

**Mechanism of action:** blocks presynaptic alpha 2 and 5HT$_2$ and 3 receptors increasing central noradrenaline and serotonin (blocking 5HT$_2$ and 5HT$_3$ receptors allowing stimulation of 5HT$_1$ receptors)

**Peak concentrations:** oral 2 hours

**Half life:** 20 to 40 hours
MORPHINE

Class: analgesic – opioid

Indications (NB some may be unlicensed): step 3 on the WHO ladder for severe pain, more effective in nociceptive than in neuropathic/visceral pains, severe breathlessness, cough, diarrhoea

Contraindications/cautions: morphine hypersensitivity/allergy (not nausea/hallucination with opioids)

Adverse reactions: common nausea/vomiting in 10 to 30% of patients (usually transient for 1 to 5 days) – give haloperidol, constipation in 90% of patients – give a stimulant & softer laxative prophylactically, dry mouth, dizziness, sedation (usually transient and on initiation or dose increase); less common respiratory depression (high doses) – pain is an antidote – give naloxone if severe, visual problems – may see things upside down/flipping, myoclonic jerking – sign of toxicity – try a different opioid, delirium in 2% of patients – give haloperidol; rare hallucinations, hyperalgesia, raised intracranial pressure, biliary/urinary tract spasm, muscle rigidity, pruritus, pulmonary oedema, physical dependence (irrelevant in dying)

Metabolism/clearance: metabolised mainly in the liver by glucuronidation to active metabolites one of which is excreted by the kidneys so watch for accumulation in renal dysfunction

Interactions:
- additive CNS effects with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), tricyclic antidepressants (e.g. amitriptyline), other opioids
- faster onset of action of slow release morphine may occur with metoclopramide

Dosing:

**pain** (initially use the normal release and titrate to pain)

oral: normal release initially 5 to 10 mg 4 hourly and prn
slow release initially 10 to 30 mg 12 hourly
- prescribe rescue doses (normal release) of 1/5th to 1/6th of the total 24 hour dose 4 to 6 hourly
- there is no real maximum dose but it is usually less than 200 mg/24 hours. If it is
- >400 mg/24 hours consider the aetiology of the pain and the use of co-analgesia or opioid rotation
- review doses regularly

subcut: oral: subcut = 2:1
rectal: oral: rectal = 1:1
epidural: subcut:epidural = 10:1
intrathecal subcut:intrathecal = 100:1

**breathlessness, cough**
oral: normal release 5 to 10 mg 4 hourly prn

**Syringe driver:** see syringe driver compatibility table

**Mechanism of action:** stimulates mu (and other) opioid receptors in the CNS and gastrointestinal tract
**Peak effect:**
- **oral:** normal release 1 hour

**Duration:**
- **oral:** normal release 4 to 5 hours
- **oral:** slow release 8 to 12 hours

**Notes:**
- Tolerance to effect does occur but progressive disease is also a cause of dose fade.
- If dose of slow release morphine is increased remember to also increase the prescribed dose of normal release morphine for breakthrough pain/rescue.
- Toxicity: decrease in respiratory rate, mental status and blood pressure – give naloxone (see naloxone page).
- For conversion to oxycodone, hydromorphone*, fentanyl or methadone, see relevant pages.
- Morphine can affect the ability to drive. Some patients may need to be told not to drive while taking morphine. Always advise patients not to drive for several days after a dose increase.
- Topical morphine may be useful for wound pain. It is usually used as 0.05 to 0.1% morphine (i.e. 0.5 to 1 mg/mL) in Intrasite™ gel, metronidazole gel or KY Jelly™.

Drugs that are either not available or not funded in New Zealand are marked with *
**MOVICOL™, LAX-SACHETS™**  
(Macrogol 3350, sodium chloride, sodium bicarbonate, potassium chloride, potassium acesulfame)  

**Class:** laxative – osmotic  
**Indications:** constipation including faecal impaction  
**Contraindications/cautions:** intestinal obstruction or perforation, ileus and severe inflammatory conditions, cardiac disease (contains sodium and potassium)  
**Adverse reactions:** less common abdominal distension and pain, nausea  
**Metabolism/clearance:** not absorbed  
**Interactions:** few as not absorbed – may affect the absorption of some drugs  
**Dosing:**  
*Movicol™, Lax-sachet™:*  
constipation 1 to 3 sachets per day  
faecal impaction 8 sachets per day taken within 6 hours for a max. of 3 days. If cardiovascular problems, do not take more than 2 sachets over any one hour  
Each sachet should be dissolved in 125 mL. For faecal impaction dissolve 8 sachets in 1 L of water  

*Movicol-Half™:*  
constipation 1 to 6 sachets/day  
faecal impaction 16 sachets/day taken within 6 hours for a max. of 3 days. If cardiovascular problems, do not take more than 4 sachets over any one hour  
Each sachet should be dissolved in 60 mL of water  

**Mechanism of action:** osmotic action in the gut to increase liquid content of stools but with no net loss of sodium, potassium or water  
**Onset:** faecal impaction most cleared after 3 days  
**Notes:**  
- Effective laxative in palliative care.  
- More acceptable to many than lactulose.
NALOXONE

Class: opioid antagonist

Indications: opioid overdose

Unlicensed indications: may enhance opioid analgesia at very low dose, may attenuate opioid adverse effects e.g. nausea and vomiting at low dose

Contraindications/cautions: cardiovascular disease

Adverse reactions: common nausea, vomiting, tachycardia, sweating, raised blood pressure (opioid withdrawal)

Metabolism/clearance: metabolised mainly in the liver by glucuronidation

Interactions:
- blocks the actions of opioids e.g. morphine, fentanyl, methadone, oxycodone, hydromorphone*

Dosing:
If respiratory rate < 8 per minute, patient unconscious or cyanosed
iv: 0.1 to 0.2 mg every 2 to 3 minutes for reversal of CNS depression post-op
  0.4 to 2 mg every 2 to 3 minutes up to 10 mg for opioid overdose
oral: not available alone
subcut: see below
rectal: not available

Syringe driver: not applicable

Mechanism of Action: blocks action of opioids at opioid receptors

Onset: iv 2 to 3 minutes subcut/im 15 minutes
Duration: 15 to 90 minutes

Notes:
- Best given iv, however if not practical can be given im or subcut.
- Reversal of respiratory depression will result in reversal of analgesia and withdrawal symptoms if physiologically dependent.
NAPROXEN

Class: non-steroidal anti-inflammatory drug (NSAID)

Indications (NB some may be unlicensed): pain associated with inflammation (including bone pain), dysmenorrhoea, itch, sweating

Contraindications/cautions: GI ulceration, asthma (in sensitive patients), renal, cardiac or hepatic impairment

Adverse reactions: common GI ulceration (more common if elderly, on steroids or aspirin), diarrhoea, indigestion, nausea; less common dizziness, rash, nephrotoxicity, hepatitis, oedema, hypertension, headache, tinnitus, proctitis (rectal administration). NB Inhibits platelet aggregation – may prolong bleeding time

Metabolism/clearance: metabolised by metabolising enzyme CYP2C8/9 mainly in the liver

Interactions:

- increased clinical effect/toxicity of naproxen (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. fluconazole, ketoconazole, voriconazole
- decreased clinical effect/toxicity of naproxen (due to decreased blood concentrations) may occur with some CYP metabolism enzyme inducers (see above) e.g. phenobarbitone, phenytoin, rifampicin
- increased clinical effect/toxicity of lithium, digoxin, methotrexate and warfarin may occur with naproxen due to increased concentrations of these drugs via kidney excretion competition so monitor
- decreased clinical effects of diuretics (e.g. frusemide) and beta blockers (e.g. propranolol) may occur with naproxen
- increased risk of renal toxicity and hyperkalaemia with ACE inhibitors (e.g. enalapril) may occur with naproxen
- additive risk of bleeding may occur with warfarin and heparin in combination with naproxen

Dosing:
oral: normal release 500 to 1,000 mg per day in two divided doses or 275 mg every 6 to 8 hours (max 1,375 mg)
sustained release 750 to 1,000 mg per day as a single dose
subcut: not available
rectal: not available (try diclofenac)

Syringe driver: not available

Mechanism of action: inhibits prostaglandin synthesis which are involved in inflammation and pain

Peak effect: oral normal release 2 to 4 hours
Duration: 7 hours
NORTRIPTYLINE

**Class:** antidepressant – tricyclic

**Indications (NB some may be unlicensed):** depression, smoking cessation, neuropathic pain, itch

**Contraindications/cautions:** arrhythmias, recent MI, epilepsy (lowers seizure threshold), urinary retention

**Adverse reactions:** _common_ anticholinergic – dry mouth, blurred vision, urinary retention, drowsiness (tolerance to these may develop except dry mouth); _less common_ sweating, constipation, confusion, arrhythmias, tachycardia, postural hypotension

**Metabolism/clearance:** metabolised by the metabolising enzyme CYP2D6 (major) mainly in the liver to active metabolites

**Interactions:**
- _increased clinical effect/toxicity of nortriptyline_ (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. bupropion, fluoxetine, paroxetine, quinine
- _additive risk of serotonin syndrome_ (potentially fatal syndrome – symptoms include sweating, diarrhoea, confusion) with other serotonergic drugs e.g. carbamazepine, fluoxetine
- _additive drowsiness_ may occur with alcohol, benzodiazepines (e.g. clonazepam)
- _increased risk of seizures in epileptics_ may occur with nortriptyline so interacts with anticonvulsants e.g. phenytoin
- _additive CNS effects_ with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), opioids, alcohol
- _additive increased risk of QT interval prolongation_ (cardiac adverse effect which may lead to arrhythmias) with other drugs that prolong the QT interval e.g. lignocaine, lithium, haloperidol

**Dosing:**

<table>
<thead>
<tr>
<th>Route</th>
<th>Depression</th>
<th>Pain</th>
</tr>
</thead>
<tbody>
<tr>
<td>oral</td>
<td>25 to 100 mg at night (max. of 50 mg in elderly)</td>
<td>10 to 50 mg at night</td>
</tr>
<tr>
<td>subcut</td>
<td>not available</td>
<td>not available</td>
</tr>
<tr>
<td>rectal</td>
<td>not available</td>
<td>not available</td>
</tr>
</tbody>
</table>

**Syringe driver:** not available

**Mechanism of action:** not really understood but thought to be through noradrenaline and serotonin in the CNS

**Onset:** depression 2 to 6 weeks  pain several days

**Notes:**
- Metabolite of amitriptyline, less adverse reactions (including sedation) than amitriptyline.
- 25 mg nortriptyline = 75 mg amitriptyline (approx).
- Measurement of blood drug concentrations may be useful to establish compliance or confirm toxicity.
NYSTATIN

Class: antifungal – polyene

Indications: fungal infections – topical, oral, gastrointestinal, vaginal

Adverse reactions: less common nausea, vomiting, diarrhoea (at high doses), local irritation

Dosing:
oral: (not absorbed orally)
gastrointestinal candidiasis: 100,000 units (1 mL) four times a day
subcut: not available
rectal: not available
topical: apply two to three times a day
vaginal: 5 g of cream once or twice a day

Syringe driver: not available

Mechanism of action: increases fungal cell membrane permeability

Notes:
• If infection is severe or recurrent use a systemic antifungal e.g. fluconazole.
**OCTREOTIDE**

**Class:** growth hormone inhibitor

**Indications NB some may be unlicensed:** acromegaly, gastro-entero pancreatic endocrine tumours, post pancreatic surgery, emergency treatment to stop bleeding oesophageal varices, antisecretory in intestinal obstruction, secretory diarrhoea, high fistula output, variceal bleeds

**Contraindications/cautions:** diabetes

**Adverse reactions:** less common injection site reaction, gastro upset, hepatitis, gallstones, hyper/hypoglycaemia, bradycardia, dizziness, drowsiness, headache, hypothyroidism

**Metabolism/clearance:** metabolised by the liver

**Interactions:**
- decreased absorption of ciclosporin may occur with octreotide

**Dosing:**
- oral: not available
- sc: 200 to 600 micrograms/24 hours (max. 1 mg/24 hours)
  - LAR – not usually used in palliative care
- rectal: not available
- iv: not available

**Syringe driver:** see syringe driver compatibility table

**Mechanism of Action:** blocks somatostatin receptors

**Peak effect:** 30 minutes

**Duration:** 12 hours

**Notes:**
- Long acting octreotide formulations are available. Their use in palliative care has not been established.
OLANZAPINE

Class: antipsychotic, antimanic, mood stabiliser

Indications (NB some may be unlicensed): acute and chronic psychoses including schizophrenia, bipolar disorder, nausea and vomiting, delirium

Contraindications/cautions: liver dysfunction, cardiovascular and cerebrovascular disease, hypotension, seizures, blood disorders, renal dysfunction, prostatic hypertrophy, paralytic ileus, bone marrow depression, diabetes, narrow angle glaucoma, hypercholesterolaemia, Parkinson’s disease, DLB

Adverse reactions: common drowsiness, weight gain, dizziness, hallucinations, akathisia and other extrapyramidal side effects, elevated blood glucose and triglycerides, chest pain, oedema, constipation, dry mouth; less common angioedema, urticaria, diabetic coma, hepatitis, pancreatitis, priapism, tardive dyskinesia, neuroleptic malignant syndrome, blood disorders, hypotension, mania, seizures

Metabolism/clearance: metabolised mainly in the liver by the metabolising enzymes CYP1A2 to inactive metabolites which are partially excreted by the kidneys

Interactions:

- increased clinical effect/toxicity of olanzapine (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. ciprofloxacin, ketoconazole
- decreased clinical effect/toxicity of olanzapine (due to decreased blood concentrations) may occur with some CYP metabolism enzyme inducers (see above) e.g. broccoli-like vegetables, smoking, phenobarbitone, phenytoin, rifampicin
- possible increase risk of extrapyramidal effects with dopamine antagonists e.g. metoclopramide
- additive hypotension with antihypertensives e.g. propranolol
- additive CNS effects with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), opioids, alcohol

Dosing:
oral tabs/disp tabs: 2.5 to 20 mg per day as a single dose
subcut: inj available but recommended for im use only
rectal: not available

Syringe driver: not available

Mechanism of action: antagonises serotonin and dopamine receptors in the CNS

Notes:
- Lower potential for neurological adverse effects than conventional antipsychotics.
- Can be used in acute delirium and behavioural disturbances associated with brain tumours.
OMEPRAZOLE

Class: ulcer healing/prophylactic – proton pump inhibitor

Indications (NB some may be unlicensed): duodenal/gastric ulcer, reflux oesophagitis, dyspepsia, NSAID associated gastric and duodenal ulcer/erosion treatment

Contraindications/cautions: renal impairment

Adverse reactions: common headache, nausea/vomiting, diarrhoea or constipation; less common insomnia, dizziness, vertigo, pruritus, blood disorders, muscle/joint pain, dry mouth, agitation

Metabolism/clearance: metabolised by metabolising enzyme CYP2C19 mainly in the liver

Interactions:
- increased clinical effect/toxicity of omeprazole (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. fluconazole, fluoxetine, ketoconazole
- decreased clinical effect/toxicity of omeprazole (due to decreased blood concentrations) may occur with some CYP metabolism enzyme inducers (see above) e.g. carbamazepine, phenytoin, rifampicin
- increased clinical effect/toxicity of some drugs (due to increased blood concentrations of them) may occur with omeprazole due to metabolising enzyme inhibition by omeprazole e.g. diazepam
- decreased absorption of itraconazole may occur with omeprazole

Dosing:
oral: 10 to 40 mg once a day
subcut: injection and infusion available but not usually used subcut. Doses of 40 mg in 100 mL normal saline have been given subcut over 3 hours
rectal: not available

Syringe driver: short infusions only

Mechanism of action: inhibits gastric acid secretion via proton pump blockade

Onset: oral (antacid effect) 10 to 20 minutes

Notes:
- Omeprazole is considered the drug of choice for prophylaxis or treatment of NSAID-induced gastro-intestinal damage.
- Oral suspension can be made.
ONDANSETRON

Class: antiemetic – 5HT₃ antagonist

Indications (NB some may be unlicensed): nausea/vomiting post chemo- or radio- therapy, post-operative nausea/vomiting, nausea/vomiting not due to above

Contraindications/cautions: hepatic impairment, subacute gastro-intestinal obstruction

Adverse reactions: common headache, constipation; less common hiccups, injection site reaction, dizziness, cardiac effects (iv usually tachycardia, chest pain, arrhythmias), sedation, convulsions

Metabolism/clearance: metabolised by metabolising enzyme CYP2D6 mainly in the liver

Interactions:
- increased clinical effect/toxicity of ondansetron (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. bupropion, fluoxetine, paroxetine, quinine

Dosing:
- oral: 4 to 8 mg twice a day
- subcut: not usually used
- rectal: not available

Syringe driver: compatibility unknown so don’t mix

Mechanism of action: acts on 5HT₃ receptors in the vomiting centre in the CNS and in the gastrointestinal tract

Peak concentration: oral 1 to 2 hours im (subcut) 30 minutes

Notes:
- May be of use in nausea and vomiting refractory to all other antiemetics.
OXYCODONE

Class: analgesic – opioid

Indications: step 3 in the WHO analgesic ladder

Contraindications/cautions: severe renal failure, respiratory disease

Adverse reactions: see morphine

Metabolism/clearance: metabolised by metabolising enzymes CYP2D6 mainly in the liver

Interactions:
- increased clinical effect/toxicity of oxycodone (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. bupropion, fluoxetine, paroxetine, quinine
- additive CNS effects with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), tricyclic antidepressants (e.g. amitriptyline), other opioids, alcohol
- additive respiratory depression with benzodiazepines (e.g. midazolam), other respiratory depressants

Dosing: (and see notes)
oral: immediate release initially in opioid naïve 1 to 3 mg 4 to 6 hourly
slow release initially 5 mg every 12 hours
subcut: oral: subcut 2:1
rectal: not available

Syringe driver: see syringe driver compatibility table

Mechanism of action: stimulates opioid receptors in the CNS and gastrointestinal tract

Onset: oral: 20 to 30 minutes

Duration: oral (immediate release) 4 to 6 hours slow release 12 hours

Notes:
- May be useful in opioid rotation.
- Dose conversion from oral morphine to oral oxycodone is 2:1 i.e. 10 mg oral morphine = 5 mg oral oxycodone because oral availability of oxycodone is twice that of morphine.
- The slow release tabs and the immediate release caps should not be opened or crushed/chewed.
- In renally impaired patients, one of oxycodone’s active metabolite may accumulate.
- The combination oxycodone+naloxone modified release tablets* are designed to reduce opioid induced constipation.

Drugs that are either not available or not funded in New Zealand are marked with *
PAMIDRONATE DISODIUM

Class: bisphosphonate calcium regulator
Indications: hypercalcaemia, metastatic bone pain, Paget’s disease
Contraindications/cautions: severe renal impairment, dental surgery, oral disease, ensure adequate hydration
Adverse reactions: less common transient flu-like symptoms, slight increase in temperature, fever, hypocalcaemia, transient bone pain, nausea, headache, osteonecrosis (particularly of jaw)
Metabolism/clearance: not metabolised, excreted by the kidneys after uptake into the bone
Interactions:
• incompatible with calcium containing infusion fluids
Dosing:
oral: not available
subcut: zoledronic acid is usually used instead
rectal: not available
iv infusion: bone pain 90 mg every 3 to 4 weeks
hypercalcaemia 15 to 90 mg depending on corrected calcium concentration
• rate of infusion should not exceed 60 mg/hour (20 mg/hour in renal impairment) and concentration should not exceed 90 mg/250 mL
Syringe driver: not applicable
Mechanism of action: inhibits bone resorption
Onset: hypercalcaemia 1 to 2 days
Duration:
hypercalcaemia 2 weeks to 3 months
bone pain 3 to 4 weeks
Notes:
• 50% of patients with metastatic bone pain may be responsive.
PANTOPRAZOLE

Class: ulcer healing/prophylaxis – proton pump inhibitor

Indications: duodenal/gastric ulcer, reflux oesophagitis, dyspepsia

Contraindications/cautions: renal impairment

Adverse reactions: common headache, nausea/vomiting; less common abdominal pain, flatulence, insomnia, pruritus, dizziness

Metabolism/clearance: metabolised by metabolising enzyme CYP2C19 mainly in the liver

Interactions:

- increased clinical effect/toxicity of pantoprazole (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. fluconazole, fluoxetine, ketoconazole
- decreased clinical effect/toxicity of pantoprazole (due to decreased blood concentrations) may occur with some CYP metabolism enzyme inducers (see above) e.g. carbamazepine, phenytoin, rifampicin
- decreased absorption of ketoconazole, itraconazole may occur with pantoprazole

Dosing:

oral: 20 to 80 mg once a day
subcut: inj available but not usually used subcut
rectal: not available

Syringe driver: not usually used

Mechanism of action: inhibits gastric acid secretion via proton pump blockade

Onset: oral (antacid effect) 2 hours
PARACETAMOL

Class: analgesic – non-opioid

Indications: step 1 on the WHO analgesic ladder, co-analgesic, antipyretic

Contraindications/cautions: severe hepatic impairment

Adverse reactions: less common rash, pancreatitis on prolonged use, liver damage in overdose (> 6 g in 24 hours) or in combination with heavy alcohol intake, nephrotoxicity

Metabolism/clearance: metabolised in the liver mainly by glucuronidation

Interactions:
- increased toxicity of paracetamol may occur with alcohol
- increased anticoagulant effect of warfarin may occur if given with concurrent paracetamol regularly for a long time so monitor INR
- increased absorption of paracetamol may occur with metoclopramide and domperidone
- increased risk of hepatotoxicity may occur with concurrent carbamazepine, phenytoin

Dosing:
oral: 500 mg to 1 g 4 to 6 hourly (max. 4 g in 24 hours)
subcut: infusion available but large volume
rectal: as for oral

Syringe driver: not used subcut due to high volume

Mechanism of Action: thought to have a central effect on pain pathways and not anti-inflammatory

Onset: 0.5 hours

Duration: 4 hours

Notes:
- Give regularly rather than if required.
- Combination preparations are not recommended.
- Liver damage is likely to occur in overdose.
- Useful analgesic when given regularly in combination with opioids.
PHENOBARBITONE

Class: anticonvulsant – barbiturate

Indications (NB some may be unlicensed): seizure control, status epilepticus, pre-op anxiety, terminal restlessness

Contraindications/cautions: acute intermittent porphyria, elderly, renal/hepatic failure

Adverse reactions: common drowsiness, headache; less common GI upset, paradoxical excitement, pain, hypocalcaemia

Metabolism/clearance: may be metabolised by metabolising enzyme CYP2C19 mainly in the liver

Interactions:
- increased clinical effect/toxicity of phenobarbitone (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. fluconazole, fluoxetine, ketoconazole
- decreased clinical effect/toxicity of phenobarbitone (due to decreased blood concentrations) may occur with some CYP metabolism enzyme inducers (see above) e.g. phenytoin, rifampicin
- decreased clinical effect/toxicity of some drugs (due to decreased blood concentrations of them) may occur with phenobarbitone due to metabolising enzyme induction by phenobarbitone e.g. aprepitant, buspirone, carbamazepine, clonazepam, dexamethasone, diazepam, domperidone, fentanyl, itraconazole, ketoconazole, methadone, midazolam, NSAIDs (e.g. diclofenac), phenytoin, prednisone, quetiapine, triazolam, warfarin
- additive CNS effects with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), opioids, alcohol

Dosing: terminal agitation
oral: 60 to 180 mg per day
subcut: 600 to 1,200 mg/ 24 hours
rectal: not available

Syringe driver: give alone and watch for irritation at injection site

Mechanism of action: depresses activity of all excitable tissue perhaps via GABA

Notes:
- Risk of respiratory depression in overdose.
PHENYTOIN

Class: anticonvulsant – hydantoin

Indications (NB some may be unlicensed): epilepsy, prophylaxis in neurosurgery, arrhythmias

Contraindications/cautions: low albumin

Adverse reactions: common gingival hyperplasia; less common slurred speech, confusion, dizziness, blood disorders, skin reactions, hepatitis

Metabolism/clearance: metabolised by metabolising enzyme CYP2C8/9 mainly in the liver

Interactions:
- increased clinical effect/toxicity of phenytoin (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. fluconazole, ketoconazole, voriconazole
- decreased clinical effect/toxicity of phenytoin (due to decreased blood concentrations) may occur with some CYP metabolism enzyme inducers (see above) e.g. phenobarbitone, rifampicin
- decreased clinical effect/toxicity of some drugs (due to decreased blood concentrations of them) may occur with phenytoin due to metabolising enzyme induction by phenytoin e.g. aprepitant, buspirone, amitriptyline, carbamazepine, clonazepam, dexamethasone, diazepam, domperidone, fentanyl, itraconazole, ketoconazole, methadone, midazolam, NSAIDs (e.g. diclofenac), olanzapine, ondansetron, phenytoin, prednisone, quetiapine, triazolam, warfarin

Dosing:
oral: 100 to 300 mg/24 hours (titrate to plasma concentrations)
subcut: inj available but not given subcut
rectal: not available

Syringe driver: not applicable

Mechanism of action: inhibits spread of seizure through the motor cortex possibly via sodium channels

Peak response: 7 to 10 days (if loaded 8 to 12 hours)

Notes:
- Monitor plasma concentrations.
- Small dose increases may result in large plasma concentration increases.
- If the patient has NG feeds these will affect phenytoin concentrations.
PREDNISONE

Class: corticosteroid – glucocorticoid

Indications (NB some may be unlicensed): allergy, asthma, rheumatic disease, inflammatory conditions, nausea/vomiting, inflammation in gastrointestinal obstruction, sweating, itch, hypercalcaemia, hiccup, pain, dyspnoea (lymphangitis), liver capsule pain, tenesmus

Contraindications/cautions: infections, gastrointestinal bleeding, diabetes, congestive heart failure, mood disorders

Adverse reactions: common insomnia (decrease by giving as single dose in the morning); less common sodium/fluid retention, GI ulceration, delayed wound healing, thinning of skin (on prolonged use), proximal muscle weakness, Cushing’s syndrome, weight gain, depression, mania, delirium

Metabolism/clearance: metabolised by the metabolising enzyme CYP3A mainly in the liver

Interactions:
- increased clinical effect/toxicity of prednisone (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. aprepitant, clarithromycin, fluconazole, fluoxetine, grapefruit juice, irtraconazole, ketoconazole, valproate
- decreased clinical effect/toxicity of prednisone (due to decreased blood concentrations) may occur with some CYP metabolism enzyme inducers (see above) e.g. carbamazepine, phenobarbitone, phenytoin, rifampicin, St John’s wort
- increased risk of GI bleed/ulceration when given with NSAIDs (e.g. diclofenac)

Dosing:
- oral: 10 to 100 mg usually once a day (max. 250 mg/day)
- subcut: not available
- rectal: not available

Syringe driver: not available

Mechanism of action: decreases inflammatory response thought to be via induction of lipocortin, an anti-inflammatory protein

Notes:
- 0.75 mg dexamethasone has an equivalent anti-inflammatory effect to 5 mg prednisone or
- 20 mg hydrocortisone.
- On discontinuation decrease dose slowly (taper) unless the patient has been taking it for less than five days in which case dose tapering is not necessary.
- Alteration in mood not usually seen below 40 mg prednisone (6 mg dexamethasone) per day.
- Corticosteroid induced insomnia responds to benzodiazepines (e.g. temazepam).
- Corticosteroid induced mood disorder is usually depression and rarely mania.
- Metabolised to prednisolone.
PREGABALIN*

Class: anticonvulsant

Indications (NB some may be unlicensed): neuropathic pain, adjunctive anticonvulsant

Contraindications/cautions: renal disease (reduce dose)

Adverse reactions: common dizziness, somnolence, blurred vision, fatigue, dry mouth, headache, tremor, constipation, nausea; less common weight gain, ataxia, confusion, suicidal thoughts

Metabolism/clearance: not metabolised, mainly excreted unchanged by the kidneys (fraction excreted unchanged by the kidney = 0.9) so adjust dose in renal dysfunction

Interactions:
  • additive CNS depression with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), tricyclic antidepressants (e.g. amitriptyline), opioids, alcohol

Dosing:
oral: neuropathic pain/epilepsy 150 to 600 mg/day in two divided doses (start with lower dose and increase)
subcut: not available
rectal: not available

Syringe driver: not available

Mechanism of action: may act through effects on calcium channels in the CNS and reduces release of the neurotransmitters glutamate, noradrenaline and substance P.
QUETIAPINE

Class: antipsychotic – atypical

Indications (NB some may be unlicensed): acute and chronic psychoses including schizophrenia, manic episodes associated with bipolar disorder, nausea and vomiting, delirium

Contraindications/cautions: liver dysfunction, cardiovascular and cerebrovascular disease, hypotension, seizures, Parkinsons, DLB

Adverse reactions: common drowsiness, dry mouth, GI effects, tachycardia, dizziness, headache, agitation, insomnia, weight gain, dyspepsia; less common neuroleptic malignant syndrome, tardive dyskinesia, cholesterol changes, thyroid hormone changes, peripheral oedema, diabetes, extrapyramidal adverse effects, hepatotoxicity, blood disorders, postural hypotension, seizures, dyspnoea, sweating, rash

Metabolism/clearance: metabolised almost completely mainly in the liver by the metabolising enzyme CYP3A

Interactions:

- increased clinical effect/toxicity of quetiapine (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. aprepitant, clarithromycin, grapefruit juice, indinavir, itraconazole, ketoconazole, nelfinavir, ritonavir, telaprevir, voriconazole
- decreased clinical effect/toxicity of quetiapine (due to decreased blood concentrations) may occur with some CYP metabolising enzyme inducers (see above) e.g. carbamazepine, phenobarbitone, phenytoin, rifampacin, St John’s wort
- possible increase risk of extrapyramidal effects with dopamine antagonists e.g. metoclopramide
- additive hypotension with antihypertensives e.g. propranolol may occur
- additive CNS effects with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), opioids, alcohol

Dosing:

oral: psychosis initially 50 mg/day increasing daily to 150 to 750 mg per day in 2 divided doses

mania initially 100 mg/day increasing daily to 200 to 800 mg per day in 2 divided doses

tranquilisation, sedation, antiemetic 25 to 100 mg at night

subcut: not available

rectal: not available

Syringe driver: not available

Mechanism of action: antagonises serotonin and dopamine receptors in the CNS

Notes:

- Lower potential for neurological adverse effects (e.g. extrapyramidal effects) than conventional antipsychotics.
- Can be used in acute delirium and behavioural disturbances associated with brain tumours.
RANITIDINE

Class: ulcer healing/prophylactic – H₂ antagonist

Indications (NB some may be unlicensed): duodenal/gastric ulcer, reflux oesophagitis, dyspepsia, itch, sweating

Contraindications/cautions: renal impairment

Adverse reactions: common diarrhoea, tiredness; less common blurred vision, gynaecomastia, bradycardia, tachycardia, hypotension, agitation, hallucinations, blood disorders, dizziness, headache, confusion

Metabolism/clearance: metabolised by the liver to 3 inactive metabolites which are excreted by the kidney together with 30% of the parent drug

Interactions:
- increased anticoagulation effect of warfarin may occur
- decreased absorption of itraconazole, ketoconazole may occur
- increased clinical effect/toxicity of metformin, oral midazolam may occur

Dosing:
- oral: 150 mg twice a day or 300 mg at night (reduce dose in elderly and renal impairment)
- subcut: 100 to 200 mg/24 hours
- rectal: not available

Syringe driver: ? infuse alone

Mechanism of Action: inhibits gastric acid secretion via histamine receptor blockade

Onset (acid suppression): oral 10 to 20 minutes

Notes:
- Pantoprazole or omeprazole is considered the drug of choice for prophylaxis or treatment of NSAID-induced gastrointestinal damage.
- If gastrointestinal reflux is uncontrolled by pantoprazole or omeprazole, adding in a night-time dose of ranitidine may help.
RISPERIDONE

Class: antipsychotic – atypical

Indications (NB some may be unlicensed): schizophrenia, psychosis, behavioural/psychological symptoms of dementia, conduct/behavioural disorders in mentally retarded, autism, mania in bipolar disorder, delirium

Contraindications/cautions: Parkinson’s disease, DLB, epilepsy, cardiovascular/cerebrovascular disease, diabetes

Adverse reactions: common insomnia, anxiety, headache, extrapyramidal symptoms; less common drowsiness, dizziness, GI upset, sexual dysfunction, constipation, dry mouth, postural hypotension

Metabolism/clearance: metabolised by metabolising enzyme CYP3A and 2D6 mainly in the liver

Interactions:
• increased clinical effect/toxicity of risperidone (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. aprepitant, bupropion, clarithromycin, fluoxetine, grapefruit juice, indinavir, itraconazole, ketoconazole, nelfinavir, paroxetine, quinine, ritonavir, telaprevir, voriconazole
• decreased clinical effect/toxicity of prednisone (due to decreased blood concentrations) may occur with some CYP metabolism enzyme inducers (see above) e.g. carbamazepine, phenobarbitone, phenytoin, rifampicin, St John’s wort
• possible increased risk of extrapyramidal effects with dopamine antagonists e.g. metoclopramide
• additive hypotension may occur with antihypertensives e.g. enalapril
• additive CNS effects with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), opioids, alcohol

Dosing:
oral: schizophrenia initially 2 mg/day increasing to 4 to 6 mg/day (max 16 mg/day)
bipolar mania initially 2 mg/day increasing to 2 to 6 mg/day dementia initially 0.25 mg twice a day increasing to a max. of 1 mg twice a day
psychosis 0.5 to 4 mg twice a day

subcut/rectal: not available

Syringe driver: not available

Mechanism of Action: antagonises serotonin and dopamine receptors in the CNS

Onset: psychosis 1 to 2 weeks

Notes:
• Lower potential for neurological adverse effects e.g. extrapyramidal effects than conventional antipsychotics.
• Increasingly used in acute delirium and behavioural disturbances associated with brain tumours.
• At high dose (> 6 to 8 mg a day) or in the cerebrally compromised patient extrapyramidal side effects may occur.
SENNA

Class: laxative – stimulant

Indications: constipation

Contraindications/cautions: acute abdominal pain, intestinal obstruction

Adverse reactions: common abdominal cramps, diarrhoea, perianal irritation; less common atonic colon (with prolonged use), hypokalaemia, discolouration of urine (brown or pink)

Metabolism/clearance: not absorbed to a great extent

Interactions:
• decreased antispasmodic effects of antispasmodics e.g. hyoscine butylbromide may occur

Dosing:
oral: 2 to 4 tabs (14 to 28 mg) at night with docusate 1 to 2 tabs at night (max. 4 tabs)
subcut: not available
rectal: not available

Syringe driver: not available

Mechanism of Action: stimulates colonic activity via nerves in the intestinal mucosa. May also have stool softening properties.

Onset: 6 to 12 hours

Notes:
• May be useful in opioid induced constipation.
SPIRONOLACTONE

Class: diuretic – aldosterone antagonist, potassium sparing

Indications (NB some may be unlicensed): oedema, hypertension, congestive heart failure, hirsutism, hyperaldosteronism, malignant ascites

Contraindications/cautions: moderate/severe renal dysfunction, hyperkalaemia, hyponatraemia

Adverse reactions: common GI upset, drowsiness, hyperkalaemia; less common rashes, headache, confusion, impotence, gynaecomastia, hyponatraemia

Metabolism/clearance: metabolised in liver to active metabolites which are excreted partially by the kidneys

Interactions:
- increased risk of hyperkalaemia with NSAIDs (e.g. diclofenac), ACE inhibitors (e.g. cilazapril, quinapril), potassium supplements
- increased clinical effect/toxicity of digoxin may occur via increased digoxin concentrations

Dosing:
oral: malignant ascites 100 to 200 mg once a day (max. 400 mg daily)
subcut/rectal: not available

Syringe driver: not available

Mechanism of Action: inhibits aldosterone causing naturesis and potassium retention

Peak response:
ardosterone antagonism 6 to 8 hours
reduced ascites 10 to 25 days

Notes:
- Paracentesis may be necessary in malignant ascites.
- Monitor body weight and renal function.
TRAMADOL

**Class:** analgesic – opioid (with extra effect on inhibitory pain pathways)

**Indications:** step 2 on the WHO analgesic ladder

**Contraindications/cautions:** epilepsy, drug abuse, respiratory depression

**Adverse reactions:** *common* nausea, vomiting, diarrhoea, sweating (dose related); *less common* dry mouth, sedation, headache, hypertension, confusion

**Metabolism/clearance:** metabolised by metabolising enzyme CYP2D6 mainly in the liver to an active metabolite

**Interactions:**
- *increased clinical effect/toxicity of tramadol* (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. bupropion, fluoxetine, paroxetine, quinine
- *additive CNS effects* with other CNS depressants e.g. benzodiazepines (e.g. lorazepam), phenothiazines (e.g. chlorpromazine), other opioids, alcohol
- *additive risk of serotonin syndrome* (potentially fatal syndrome – symptoms include sweating, diarrhoea, confusion) with other serotonergic drugs e.g. amitriptyline, carbamazepine, citalopram, fluoxetine, lithium, paroxetine
- *decreases seizure threshold* so may interact with anticonvulsants e.g. carbamazepine

**Dosing:**
- **oral:** normal release 50 to 100 mg 4 hourly (max. 400 mg/24 hours)
  - slow release 100 to 200 mg twice a day
- **subcut:** up to 600 mg/24 hours
- **rectal:** not available

**Syringe driver:** give separately as compatibility as yet unknown

**Mechanism of action:** stimulates mu opioid receptors in CNS and gastrointestinal tract and also affects noradrenaline and serotonin in descending spinal inhibitory pain pathways

**Peak effect:** oral normal release 0.5 to 1 hour

**Duration:** oral normal release 3 to 7 hours

**Notes:**
- Place in palliative therapy still to be established.
- May be useful in patients who are constipated on codeine as it is less constipating generally.
- Start with low dose to minimise adverse effects.
- It is not a controlled drug.
TRANEXAMIC ACID

Class: antifibrinolytic, haemostatic

Indications: haemorrhage – surface bleeding from tumours, nose and other organs

Contraindications/cautions: active clotting, urinary tract bleeds (as clots may rarely form in the urinary tract), renal dysfunction, subarachnoid haemorrhage, acquired defective colour vision

Adverse reactions: common Gl upset less common dizziness (iv), thrombocytopenia, headache, restlessness, impaired colour vision

Interactions:
• decreased clinical effect of anticoagulants e.g. warfarin may occur with tranexamic acid

Dosing: haemorrhage
oral: 1 to 1.5 g three to four times a day
subcut: not used
rectal: the injection has been used rectally for rectal bleeding
topical: the injection has been used topically on bleeding wounds
iv: 0.5 to 1 g two to three times a day

Syringe driver: not applicable

Mechanism of action: interacts with plasminogen to cause antifibrinolysis

Peak effect: 3 hours

Notes:
• Tablets are large and many patients may have difficulty swallowing them.
VALPROATE (SODIUM)

Class: anticonvulsant, antipsychotic

Indications (NB some may be unlicensed): epilepsy, bipolar disease, neuropathic pain

Contraindications/cautions: liver dysfunction

Adverse reactions: common GI upset, tremor; less common thrombocytopenia, sedation, transient hair loss, hepatotoxicity

Metabolism/clearance: may be metabolised by CYP metabolising enzymes family mainly in the liver

Interactions:
- increased clinical effect/toxicity of some drugs (due to increased blood concentrations of them) may occur variably with valproate due to metabolising enzyme inhibition by valproate e.g. amitriptyline, carbamazepine, citalopram, NSAIDs (e.g. diclofenac), pantoprazole, phenobarbitone, phenytoin
- decreased clinical effect/toxicity of valproate (due to decreased blood concentrations) may occur with some CYP metabolism enzyme inducers e.g. carbamazepine

Dosing:
- neuropathic pain
  - oral: 200 to 1,000 mg twice a day (max. 2,500 mg per day, start low)
  - subcut: available in injectable form, not usually used
  - rectal: not available

Syringe driver: not applicable

Mechanism of action: pain – as for carbamazepine

Peak effect: not known but peak concentrations reached in 4 to 8 hours

Notes:
- Co-analgesic often used with opioids in the treatment of neuropathic pain although gabapentin or pregabalin have become common alternatives.
- May be used in neuropathic pain when tricyclic antidepressants have failed or in combination with tricyclic antidepressants.
- When switching from carbamazepine to valproate watch for toxicity from other drugs as carbamazepine induces the metabolism of several drugs while valproate inhibits the metabolism of several drugs.
- Don’t discontinue abruptly as risk of rebound seizures.
- Therapeutic drug monitoring is usually available but is of limited value.
- Monitor LFTs.
VENLAFAXINE

Class: antidepressant – bicyclic, SNRI

Indications (NB some may be unlicensed): depression, anxiety disorders, neuropathic pain, hot flushes

Contraindications/cautions: renal/hepatic failure, volume depletion, epilepsy, mania, heart disease

Adverse reactions: common nervousness, headache, fatigue, blood pressure changes, dizziness, dry mouth, insomnia, drowsiness, weight gain or loss, GI effects, sexual dysfunction, sweating, weakness, prolongation of the QT interval; less common tremor, mania, anxiety, palpitations, heart failure, loss of consciousness, seizures, blood disorders, hepatitis, arrhythmias, neuroleptic malignant syndrome, pancreatitis, extrapyramidal adverse effects, hypercholesterolaemia

Metabolism/clearance: metabolised by metabolising enzyme CYP2D6 mainly in the liver to active metabolites. Some venlafaxine and some of its metabolites are excreted by the kidneys

Interactions:

- increased clinical effect/toxicity of venlafaxine (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. bupropion, fluoxetine, paroxetine, quinine
- increased clinical effect/toxicity of some drugs (due to increased blood concentrations of them) may occur with venlafaxine due to metabolising enzyme inhibition e.g. codeine (effect may be decreased due to lack of metabolism to morphine), nortriptyline
- increased risk of serotonin syndrome with MAOIs e.g. phenelzine so avoid venlafaxine within 2 weeks of MAOI therapy
- increased risk of prolonged QT interval with other drugs that prolong the interval e.g. haloperidol

Dosing:
oral: modified release 37.5 to 375 mg once a day
subcut: not available
rectal: not available

Syringe driver: not available

Mechanism of action: inhibits reuptake of serotonin (at high dose), noradrenaline and dopamine in the CNS

Notes:

- Effectiveness in neuropathic pain is yet to be evaluated.
WARFARIN

Class: anticoagulant

Indications: thrombotic disorders prophylaxis

Contraindications/cautions: potential haemorrhagic conditions

Adverse reactions: common bleeding; less common hair loss; rare purple toe syndrome

Metabolism/clearance: metabolised by the metabolising enzymes CYP 1A2, 2C19 and 2C9 mainly in the liver

Interactions:

- increased clinical effect/toxicity of warfarin (due to increased blood concentrations) may occur with some CYP metabolising enzyme inhibitors (see above) e.g. ciprofloxacin, fluconazole, fluoxetine, ketoconazole, pantoprazole
- decreased clinical effect/toxicity of warfarin (due to decreased blood concentrations) may occur with some CYP metabolism enzyme inducers (see above) e.g. broccoli like vegetables, carbamazepine, phenobarbitone, phenytoin, rifampicin, smoking
- increased risk of bleeding with aspirin, SSRIs (e.g. fluoxetine), NSAIDs (e.g. diclofenac)
- increased clinical effect of warfarin may occur with paracetamol
- decreased clinical effect of warfarin may occur with phytomenadione (vitamin K) and foods rich in vitamin K

NB Any changes in drug therapy should be accompanied by an INR check.

Dosing:

oral: adjusted to INR (see below)
subcut: not available
rectal: not available

Syringe driver: not available

Mechanism of action: interferes with vitamin K synthesis

Notes:

- A low molecular weight heparin e.g. enoxaparin may be better tolerated.
- Different brands are not proven to be equivalent.

<table>
<thead>
<tr>
<th>Treatment in DVT and PE</th>
<th>INR</th>
<th>Duration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre and perioperative anticoagulation</td>
<td>1.5 to 2.0</td>
<td>days</td>
</tr>
<tr>
<td>Treatment of calf DVT</td>
<td>2.0 to 3.0</td>
<td>4-6 weeks</td>
</tr>
<tr>
<td>Treatment of provoked DVT</td>
<td>2.0 to 3.0</td>
<td>12-26 weeks</td>
</tr>
<tr>
<td>Treatment of provoked PE or massive DVT</td>
<td>2.0 to 3.0</td>
<td>26-52 weeks</td>
</tr>
<tr>
<td>Treatment of unprovoked PE or DVT</td>
<td>2.0 to 3.0</td>
<td>life long</td>
</tr>
<tr>
<td>Treatment of recurrent PE or DVT*</td>
<td>3.0 to 4.0</td>
<td>life long</td>
</tr>
<tr>
<td>Atrial fibrillation</td>
<td>2.0 to 3.0</td>
<td>life long</td>
</tr>
<tr>
<td>Mechanical heart valves</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aortic valve replacement</td>
<td>2.0 to 2.5</td>
<td>life long</td>
</tr>
<tr>
<td>Mitral valve replacement</td>
<td>2.5 to 3.0</td>
<td>life long</td>
</tr>
<tr>
<td>Arterial disease</td>
<td>3.0 to 4.0</td>
<td>life long</td>
</tr>
</tbody>
</table>

*recurrence despite prothrombin ratio between 2 and 3.

Table from Management Guidelines for Common medical Conditions, 15th Edition 2013, Canterbury District Health Board

Drugs that are either not available or not funded in New Zealand are marked with *

148 | THE PALLIATIVE CARE HANDBOOK
ZOLEDRONIC ACID

Class: bisphosphonate – calcium regulator

Indications (NB some may be unlicensed): osteoporosis treatment and prevention, Paget’s disease, prevention of further fracture after hip fracture, hypercalcaemia of malignancy, bone metastases

Contraindications/cautions: renal or hepatic impairment, cardiac impairment, hypo-calcaemic, phosphataemic or magnesaemic patients, administration with diuretics and other nephrotoxic drugs

Adverse reactions: common hypotension, fatigue, fever and other flu-like symptoms, GI upset (nausea), rash, chest pain, renal toxicity; less common anxiety, insomnia, hypocalcaemia, hypophosphataemia and hypomagnesaemia, sore mouth/throat, eye irritation, conjunctivitis

Metabolism/clearance: excreted unchanged by the kidneys and not metabolised

Interactions:
- additive risk of renal toxicity with other nephrotoxic drugs e.g. frusemide, thalidomide

Dosing:
- oral: not available
- subcut: not usual but has been tried
- rectal: not available
- iv infusion: hypercalcaemia 4 mg iv infused over 15 mins
- bone met pain 4 mg iv as above every 3 to 4 weeks

Syringe driver: not applicable

Mechanism of action: inhibits bone resorption

Onset: hypercalcaemia 2 to 3 days

Duration:
- hypercalcaemia 32 to 39 days
- bone pain 4 to 6 weeks

Notes:
- Patients must be adequately hydrated before administration of zoledronic acid, especially the elderly and those on diuretics.
- Renal impairment has been noted after a single administration of the drug. Routinely check serum creatinine clearance pre-administration and cease zoledronic acid if creatinine is becoming impaired.
- Osteonecrosis of the jaw has been noted predominantly in adults receiving bisphosphonate infusions.
- Occasionally severe muscle, bone and joint pain is experienced after infusion, mostly this is relieved after stopping treatment.
SYRINGE DRIVERS
A syringe driver is a battery-operated pump which administers drugs subcutaneously—consult a specialist for information on the pump used in your area and how to use it. Many of the drugs administered via the syringe driver are not licensed for subcutaneous use and the responsibility for their use lies with the prescriber.

Indications
- severe nausea and/or vomiting
- dysphagia
- severe oral lesions
- non-absorption of oral medication
- unconscious or sedated patient

Diluent
- most drugs and drug combinations used in a syringe driver need to be made up to a certain number of millimeters or volume with a diluent
- generally water for injection is currently used
- some drugs, however must be diluted with a specified diluent e.g. levomepromazine (methotrimeprazine) in normal saline
- both water for injection and normal saline have advantages and disadvantages:
  - water for injection
    > has few ions present and therefore is less likely to cause precipitation of drugs out of solution
    > BUT may be more irritant to subcutaneous tissue
  - normal saline
    > contains ions and so is more likely to cause precipitation of drugs
    > BUT may be more like interstitial fluid and therefore less irritant to subcutaneous tissue
Compatibility

- often several drugs are combined in one syringe
- little work has been done on the compatibility of drugs in syringe drivers (see chart)
- examination of the drugs in the syringe may reveal visual incompatibility, e.g. precipitation BUT non-visual chemical reactions may be occurring leading to the inactivation of one or more of the drugs or the production of potentially toxic compounds
- only combine drugs that are absolutely essential – if there is any doubt, consultation with a drug information pharmacist will guide practice
- avoid combining more than three drugs in one syringe
- consider the use of more than one syringe driver when more than three drugs need to be given via this route or if there are concerns about compatibility

The following drugs should never be given subcutaneously
DIAZEPAM, PROCHLORPERAZINE, CHLORPROMAZINE
**SYRINGE DRIVER COMPATIBILITY TABLE**

| Compatibility of drugs for use in syringe drivers over 24 hours of subcutaneous infusions | clonazepam | cyclizine | dexamethasone | fentanyl | glycopyrrolate | haloperidol | hydromorphone | hyoscine butyl bromide (Buscopan™) | hyoscine hydrobromide | ketamine | methotrimeprazine/levomepromazine (Nozinan™) | methadone | methoclopramide | midazolam | morphine sulphate (normal strengths) | morphine tartrate (high strengths) | octreotide | ondansetron | oxycodone | phenobarbitone |
| clonazepam | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| cyclizine | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| dexamethasone | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| fentanyl | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| glycopyrrolate | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| haloperidol | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| hydromorphone | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| hyoscine butyl bromide (Buscopan™) | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| hyoscine hydrobromide | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| ketamine | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| methotrimeprazine/levomepromazine (Nozinan™) | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| methadone | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| methoclopramide | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| midazolam | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| morphine sulphate (normal strengths) | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| morphine tartrate (high strengths) | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| octreotide | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| ondansetron | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| oxycodone | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |
| phenobarbitone | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y | Y |

**Combinations that have been used:**

- **Y** = compatible
- **N** = incompatible
- **SI** = sometimes incompatible (usually at higher concentrations)
- **NA** = not usually used together
- **?** = unknown

| **Combinations** | **clonazepam** | **cyclizine** | **dexamethasone** | **fentanyl** | **glycopyrrolate** | **haloperidol** | **hydromorphone** | **hyoscine butyl bromide (Buscopan™)** | **hyoscine hydrobromide** | **ketamine** | **methotrimeprazine/levomepromazine (Nozinan™)** | **methadone** | **methoclopramide** | **midazolam** | **morphine sulphate (normal strengths)** | **morphine tartrate (high strengths)** | **octreotide** | **ondansetron** | **oxycodone** | **phenobarbitone** |
| morphine + clonazepam + cyclizine | compatible | | | | | | | | | | | | | | | | | | | | |
| morphine + cyclizine + dexamethasone | compatible | | | | | | | | | | | | | | | | | | | | |
| morphine + dexamethasone + haloperidol | compatible | | | | | | | | | | | | | | | | | | | | |
| morphine + clonazepam + dexamethasone | compatible | | | | | | | | | | | | | | | | | | | | |
| morphine + cyclizine + haloperidol | compatible | | | | | | | | | | | | | | | | | | | | |
| morphine + dexamethasone + hyoscine hydrobromide | compatible | | | | | | | | | | | | | | | | | | | | |
| morphine + clonazepam + haloperidol | compatible | | | | | | | | | | | | | | | | | | | | |
| morphine + cyclizine + hyoscine butyl bromide | compatible | | | | | | | | | | | | | | | | | | | | |
| morphine + dexamethone + metoclopramide | compatible | | | | | | | | | | | | | | | | | | | | |
| morphine + clonazepam + ketamine | compatible | | | | | | | | | | | | | | | | | | | | |
| morphine + cyclizine + metoclopramide | compatible | | | | | | | | | | | | | | | | | | | | |
| morphine + dexamethasone + midazolam | compatible | | | | | | | | | | | | | | | | | | | | |
| morphine + clonazepam + metoclopramide | compatible | | | | | | | | | | | | | | | | | | | | |

**Diluent:** water is recommended for all infusions except ketamine, octreotide, ondansetron and levomepromazine where sodium chloride 0.9% should be used although in combinations consider water.

Info from:
USEFUL RESOURCES

Cancer pain management guidelines (Australia)

International Association for the Study of Pain (IASP)
http://www.iasp-pain.org/

Caresearch – palliative care knowledge network

Palliativedrugs.com
http://www.palliativedrugs.com/

The Palliative Care Bridge

Palliative Care Matters
http://www.pallcare.info/

National Cancer Institute
http://www.cancer.gov/

Macmillan Cancer Support
http://www.macmillan.org.uk/

Spirituality
http://smhs.gwu.edu/gwish/global-network

Dying
http://www.dyingmatters.org

Advance care planning
http://advancecareplanning.org.au/
http://www.advancecareplanning.org.nz/


Palliative Care Victoria pamphlet ‘About Nutrition in Palliative Care’
FURTHER READING

BOOKS


Chochinov HM (2012) Dignity Therapy; final words for final days. Oxford; Oxford University Press


JOURNAL ARTICLES


systematic review and meta-analysis *Lancet Neurology*. 162–73


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