

Home Care Resources

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Home Care Pack Contents

Practical help to assist you to support older people at the end of life at home

Home care providers are a critical part of support for older people at the end of life. Ensuring that key palliative care steps are not missed, and processes are well documented is part of good practice.

To help you and your team to support older people living in their home, we have brought together this pack of resources.

Self-care plan for the aged care team

To care for others, you need to take care of yourself. This worksheet helps you to plan activities to maintain your balance in life.

Recognising changing needs

Forms for systematically assessing for deterioration in a person's health and identifying unmet supportive and palliative care needs.

• SPICT Tool • SPICT4ALL Tool.

Symptom control

Forms for monitoring for changes in pain status and tracking use of breakthrough medicines.

• Abbey Pain Scale • Breakthrough medicines.

Palliative care case conference

A series of practical checklists and forms to guide and document case conferences.

Supporting older people and their family

Forms to give to clients to help them keep track of emergency contacts, medicines, and to develop self-care plans for their well-being.

• My emergency contact list • My medicines list • Self-care plan for family carer.

End-of-life care

A practical list of medicines for responding to unanticipated needs in the terminal phase and a checklist for GPs to guide an at home death.

MBS remuneration for palliative care services

- MBS items for nurse practitioners
- MBS remuneration for GPs providing a planned palliative care pathway

To obtain further copies of this pack or other resource packs developed by palliAGED: Download or order printed copies at palliaged.com.au/practiceforms

Use QR code to download or order additional copies of these resources palliaged.com.au









Self-care plan: Aged care team

Caring for others can be rewarding. However, staff working in aged care look after many people who die. You might find this loss hard to accept. You might find it hard to sleep, no longer enjoy your work, or feel tired. This can affect you and your family. Self-care is what we do to maintain balance in our life.

A self-care plan based on what you like to do can help. We have suggested a few things that you could try, but what you choose will depend on you.

Workplace self-care: Activities to help you at work

Some examples

- Regular meetings with supervisors or a more experienced colleague
- Join a support group with the people you work with
- Attend training programs

My activities:

Physical self-care: Activities that help you to stay fit and healthy

Some examples

- Develop a regular sleep routine
- Aim for a healthy diet
- Take lunch breaks and go for a walk
- Get some exercise before/after work regularly

Self-care plan

Psychological self-care: Activities that help you to feel clear-headed and able to engage with workplace and personal challenges

Some examples

- Keep a reflective journal
- Seek regular meetings with supervisors or a more experienced colleague
- Turn off your email and work phone outside of work hours
- Make time to be with friends and family

My activities:

Emotional self-care: Allow yourself to safely express your emotions

Some examples

- Develop friendships that are supportive
- Write or think of three good things that you did each day
- Play a sport and have a coffee together after training
- Talk to your friends about how you are coping with work and life demands

Self-care plan

Spiritual self-care: Develop a sense of perspective beyond the day-to-day of life which can include religion, but it is not always about religion

Some examples

- Engage in reflective practices like meditation
- Go on walks to connect with nature
- Go to church/mosque/temple
- Do yoga

My activities:

Relationship self-care: Maintain healthy, supportive relationships, and ensure that you are not only connected to work people

Some examples

- Prioritise close relationships in your life e.g. with partners, family and children
- Attend the special events of your family and friends
- Arrive to work and leave on time every day



Supportive and Palliative Care Indicators Tool (SPICT™)



The SPICT[™] is used to help identify people whose health is deteriorating. Assess them for unmet supportive and palliative care needs. Plan care. Look for any general indicators of poor or deteriorating health.

- Unplanned hospital admission(s).
- Performance status is poor or deteriorating, with limited reversibility. (eg. The person stays in bed or in a chair for more than half the day.)
- Depends on others for care due to increasing physical and/or mental health problems.
- The person's carer needs more help and support.
- Progressive weight loss; remains underweight; low muscle mass.
- Persistent symptoms despite optimal treatment of underlying condition(s).
- The person (or family) asks for palliative care; chooses to reduce, stop or not have treatment; or wishes to focus on quality of life.

Look for clinical indicators of one or multiple life-limiting conditions.

Cancer

Functional ability deteriorating due to progressive cancer.

Too frail for cancer treatment or treatment is for symptom control.

Dementia/ frailty

Unable to dress, walk or eat without help.

Eating and drinking less; difficulty with swallowing.

Urinary and faecal incontinence.

Not able to communicate by speaking; little social interaction.

Frequent falls; fractured femur.

Recurrent febrile episodes or infections; aspiration pneumonia.

Neurological disease

Progressive deterioration in physical and/or cognitive function despite optimal therapy.

Speech problems with increasing difficulty communicating and/or progressive difficulty with swallowing.

Recurrent aspiration pneumonia; breathless or respiratory failure.

Persistent paralysis after stroke with significant loss of function and ongoing disability.

Heart/ vascular disease

Heart failure or extensive, untreatable coronary artery disease; with breathlessness or chest pain at rest or on minimal effort.

Severe, inoperable peripheral vascular disease.

Respiratory disease

Severe, chronic lung disease; with breathlessness at rest or on minimal effort between exacerbations.

Persistent hypoxia needing long term oxygen therapy.

Has needed ventilation for respiratory failure or ventilation is contraindicated.

Other conditions

Kidney disease

Stage 4 or 5 chronic kidney disease (eGFR < 30ml/min) with deteriorating health.

Kidney failure complicating other life limiting conditions or treatments.

Stopping or not starting dialysis.

Liver disease

Cirrhosis with one or more complications in the past year:

- diuretic resistant ascites
- hepatic encephalopathy
 - hepatorenal syndrome
 - bacterial peritonitis
 - recurrent variceal bleeds

Liver transplant is not possible.

Deteriorating with other conditions, multiple conditions and/or complications that are not reversible; any treatment available will have a poor outcome.

Review current care and care planning.

- Review current treatment and medication to ensure the person receives optimal care; minimise polypharmacy.
- Consider referral for specialist assessment if symptoms or problems are complex and difficult to manage.
- Agree a current and future care plan with the person and their family. Support carers.
- Plan ahead early if loss of decision-making capacity is likely.
- Record, share, and review care plans.

Please register on the SPICT website (www.spict.org.uk) for information and updates

SPICTTM, 2022



Supportive and Palliative Care Indicators Tool (SPICT-4ALL™)



The SPICT[™] helps us to look for people who are less well with one or more health problems. These people need more help and care now, and a plan for care in the future. Ask these questions:

Does this person have signs of poor health or health problems that are getting worse?

- Unplanned (emergency) admission(s) to hospital.
- General health is poor or getting worse; the person never quite recovers from being more unwell. (This means the person is less able to manage day to day life and often stays in bed or in a chair for more than half the day).
- Needs help from others for care due to increasing physical and/ or mental health problems.
- The person's carer needs more help and support.
- Has clearly lost weight over the last few months; or stays too thin.
- Has troublesome symptoms most of the time despite good treatment of their health problems.
- The person (or family) asks for palliative care; chooses to reduce, stop or not have treatment; or wishes to focus on quality of life.

Does this person have any of these health problems?

Cancer

Less able to manage usual activities; health getting poorer.

Not well enough for cancer treatment or treatment is to help with symptoms.

Dementia/frailty

Unable to dress, walk or eat without help.

Eating and drinking less; difficulty with swallowing.

Has lost control of bladder and bowels.

Not able to communicate by speaking; not responding much to other people.

Frequent falls; fractured hip.

Frequent infections; pneumonia.

Nervous system problems

(eg Parkinson's disease, MS, stroke, motor neurone disease)

Physical and mental health are getting worse.

More problems with speaking and communicating; swallowing is getting worse.

Chest infections or pneumonia; breathing problems.

Severe stroke with loss of movement and ongoing disability.

Heart or circulation problems

Heart failure or has bad attacks of chest pain. Short of breath when resting, moving or walking a few steps.

Very poor circulation in the legs; surgery is not possible.

Lung problems

Unwell with long term lung problems. Short of breath when resting, moving or walking a few steps even when the chest is at its best.

Needs to use oxygen for most of the day and night.

Other conditions

Has needed treatment with a breathing machine in the hospital.

Kidney problems

Kidneys not working well; general health is getting poorer.

Stopping kidney dialysis or choosing supportive care instead of starting dialysis.

Liver problems

Worsening liver problems in the past year with complications like:

- fluid building up in the belly
- being confused at times
- kidneys not working well
- infections
- bleeding from the gullet

A liver transplant is not possible.

People who are less well and may die from other health problems or complications. There is no treatment available or it will not work well.

What we can do to help this person and their family.

- Start talking with the person and their family about any help needed now and why making plans for care is important in case things change.
- Ask for help and advice from a nurse, doctor or other professional who can assess the person and their family and help plan care.
- We can look at the person's medicines and other treatments to make sure we are giving them the best care or get advice from a specialist if problems are complicated or hard to manage.
- We need to plan early if the person might not be able to decide things in the future.
- We make a record of the care plan and share it with people who need to see it.

SPICT-4ALLTM 2023

Abbey Pain Scale

Enter pain scores for each of the the following six areas:

Absent = 0; Mild = 1; Moderate = 2; Severe = 3

Patient details	
Surname	
Title	
Given names	
DOB	MRN
Address	
Suburb	
Postcode	

		T			,
Enter date:					
Enter time:					
Sign entry					
1. Vocalisation e.g., whimpering, groaning, crying.					
2. Facial expression e.g., looking tense, frowning, grimacing, looking frightened.					
3. Change in body language e.g., fidgeting, rocking, guarding part of body, withdrawn.					
4. Behavioural change e.g., increased confusion, refusing to eat, alteration in usual patterns.					
5. Physiological change e.g., temperature, pulse or blood pressure outside normal limits, perspiring, flushing or pallor.					
6. Physical changes e.g., skin tears, pressure areas, arthritis, contractures, previous injuries.					
Total scores					
Circle the range that matches the total pain score					
0-2 No pain	No pain	No pain	No pain	No pain	No pain
3-7 mild	Mild	Mild	Mild	Mild	Mild
8-13 moderate 14+ severe	Moderate	Moderate	Moderate	Moderate	Moderate
	Severe	Severe	Severe	Severe	Severe

Chronic

Tick the box which matches the type of pain: Acute

Acute on chronic

About Abbey Pain Scale

Purpose: Developed to detect pain in elderly residents with dementia and inability to communicate verbally. It is a 6-item 3 point scale tool.

Description: The Abbey Pain Scale was developed for use in aged care and dementia care. The tool is best used as part of an overall pain management plan. As the tool does not differentiate between distress and pain measuring the effectiveness of any interventions is essential. Use the form to collate recordings across an extended period to facilitate monitoring of responses. The Australian Pain Society recommends using the tool as a movement-based assessment and conducting a **second evaluation one hour after any intervention taken.** Repeat hourly until a score of mild pain is reached and then 4 hourly for 24 hours with treatment for pain as required. Contact the GP or pain team if there is no improvement.

Acknowledgement: Abbey J, et al. The Abbey pain scale: A 1-minute numerical indicator for people with end-stage dementia. Int J Palliat Nurs. 2004 Jan;10(1):6-13.

Chart of breakthrough medicines

Patient details	
Surname	
Title	
Given names	
DOB MRN	
Address	
Suburb	
Postcode	

About breakthrough medicines

Breakthrough symptoms occur for many reasons in people with palliative care needs. This includes symptoms like pain, nausea, breathlessness, and anxiety. Sometimes these symptoms 'break through' the stable control of symptoms that you might usually experience. Your prescriber may prescribe additional medicines to be used as a 'rescue dose' in case this happens. It is important that you follow the advice given to you about treating breakthrough symptoms. Letting your care team know how much and how often breakthrough medicine is needed helps with management of your symptoms. You can use this chart to keep track of any breakthrough medicine that you or the person you are caring for takes.

1. Medicine name:

Strength:

How much and how often to be given:

Reason for taking:

Date				
Time				
Dose				
Time				
Dose				
Time				
Dose				
24-hour total dose				

2. Medicine name:

Strength:

How much and how often to be given:

Reason for taking:

Date				
Time				
Dose				
Time				
Dose				
Time				
Dose				
24-hour total dose				

3. Medicine name:

Strength:

How much and how often to be given:

Reason for taking:

Date				
Time				
Dose				
Time				
Dose				
Time				
Dose				
24-hour total dose				

My Emergency Contact List:

Organisation:

Helping you keep track of your team

My name:

Reminder:

Put son fou

Put this list on your fridge or somewhere where it can be found.

Relationship/ Role	Name	Phone Number	Contact at time of death? (Y/N)
Partner/friend/ family member			
Substitute Decision-maker			
Specialist			
General Practitioner (GP)			
Nurse			
Pharmacist			
Other			

Medicines List:

Helping you keep track of your medicines

My name:

My allergies or previous problems:

My emergency contact(s) details:

My GP/specialist contact details:

My pharmacy:

My pharmacist(s):

My palliative care team (e.g., careworker, nurse):

Reminders:

- Ask a member of your care team to help you fill out this form.
- Bring this form to any future medical appointments.
- Include non-prescription medicines.

Name of medicine	What it looks like	How much and when	How to take it	Date started	What the medicine is for
Example only	e.g., round, red, blue, white liquid	e.g., one capsule per day	e.g., by mouth, with food, by injection	dd/mm/yy	e.g., pain

Self-care plan:

Family carer

Name:

Caring for someone at home at the end of life is complicated and you may find it challenging. The intensity of the caring situation can be hard to deal with. You might find it hard to sleep, feel anxious or worried. Your friends may not visit as often. This can affect you and your family. Self-care is what we do to maintain balance in our life.

A self-care plan based on what you like to do can help. Here we suggest a few things that you could try, but what you choose will depend on what suits you best.

Home self-care: Activities to help you at home

Some examples

Have someone to help with the care so that you can get time for yourself

Share an online calendar with family and friends so that they know when you might need help or company

Find out about respite services in your area that can be accessed for a few hours or a few days Join a support or training group for carers

Attend training programs for carers

My activities:

Physical self-care: Activities that help you to stay fit and healthy

Some examples

Develop a regular sleep routine

Aim for a healthy diet

Take lunch breaks and go for a walk

Get some exercise before/after work regularly

Self-care plan

Psychological self-care: Activities that help you to feel clear-headed and able to engage with personal challenges

Some examples

Keep a reflective journal

Seek regular meetings with the palliative care team

Make time to be with friends and family

My activities:

Emotional self-care: Allowing yourself to safely express your emotions

Some examples

Develop friendships that are supportive

Write or think of three good things that you did each day

Play a sport and have a coffee together after training

Talk to your friends about how you are coping with work and life demands

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Self-care plan

Spiritual self-care: Develop a sense of perspective beyond the day-to-day of life which can include religion, but it is not always about religion

Some examples

Engage in reflective practices like meditation

Go on walks to connect with nature

Go to church/mosque/temple

Do yoga

My activities:

Relationship self-care: Maintain healthy, supportive relationships

Some examples

Prioritise close relationships in your life e.g.. with partners, family and children **Attend** the special events of your family and friends

Using the palliAGED palliative care case conference forms

A case conference or family meeting between the person, their family and care providers can help to explain what is happening and to plan care. The palliAGED forms can help.

> Use the palliAGED case conference checklist for residential care or for home care to organise a palliative care case conference. Tick off items as they are completed. Speak with the person and their family about the need for a case 2 conference. Provide information on palliative care and case conferences. Involving the person's GP is important. Use the GP invitation to 3 invite them to attend, and/or to suggest a suitable time. Closer to the date of the case conference, send a letter confirming

details to the person and their family, and send confirmation to the GP.

To guide the meeting and to make sure that all steps following the conference are completed use the palliAGED <u>case conference</u> <u>summary for residential care</u> or <u>case conference summary for</u> <u>home care</u> sheet.

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Planning checklist: Home care

Organisation:

Palliative care case conference

Full name of client:	DOB (dd/mm/yy):				
Date of case conference (dd/mm/yy):	Time:				
Venue:	Room booked:				
Dial-in telephone number:	Code:				
Case conference facilitator:					
Goals of case conference:					

Family participants

Name	Role/relationship	Contact details

Health and Care Professionals

Name	Role/relationship	Contact details

Document (tick as appropriate)	Sent	Accepted/declined		N/A
Client & family information		А	D	
Client & family confirmation		А	D	
GP invitation		А	D	
GP confirmation		А	D	

	Needed	Obtained	N/A
Clinical record (including most recent medication chart)			
Advance care planning document (legal or non-legal)			
Carer document e.g. NAT-C needs assessment form			
Other (specify)			

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GP invitation

Organisation:

Palliative care case conference

То:	Email/fax number:			
From:	No. of pages: (including this page)			
Subject: Palliative case conference	Date sent: (dd/mm/yy):			
Dear Dr				
A palliative care case conference is being orga	anised for (resident/client name):			
Resident/client DOB (dd/mm/yy):				
Proposed date (dd/mm/yy):	Start time:			
Expected duration:	Venue:			
Reason for case conference:				
Please indicate availability to participate in below:	this case conference by ticking one of the options			
Attending in person	Unable to attend			
Attending via teleconference Please provide your telephone number:				
Please reschedule so I can attend. Proposed alternative date: (dd/mm/yy):	and time:			
Please email/fax this back to (insert email/fax number):				
Yours sincerely (name):				
Role:	Organisation:			

palliAGED Practice Resources

GP confirmation

Organisation:

Palliative care case conference

То:	Email/fax number:
From:	No. of pages: (including this page)
Subject: Palliative case conference	Date sent: (dd/mm/yy):
Dear Dr	
Following our recent correspondence wit for: (resident/client name):	h you a palliative care case conference has been organised
Resident/client DOB (dd/mm/yy):	
Case conference date (dd/mm/yy):	Start time:
Case conference date (dd/mm/yy):	Venue:
Expected duration:	Venue:

depression

• delirium (sudden confusion)

- nausea (feel that you want to vomit)

Role:

Information for you and your family

Palliative care case conferences

It has been suggested that a case conference be held to discuss how you, or your family member might benefit from palliative care. The following explains what this is and why it is important.

Case conference: Case conferences or family meetings are an opportunity to discuss a person's care needs. They ideally include the person (if able to attend), their family and/or their substitute decision-maker, and members of the care team including the doctor.

Palliative care: Palliative care is person- and family-centred care that supports a person to live the best life they can with a life-limiting illness. A life-limiting illness means that the person has little or no prospect of cure and is expected to die. The focus is on quality of life.

Life-limiting illnesses include dementia, advanced heart, kidney, lung or liver disease, cancer, and motor neurone disease.

People can receive palliative care for days or weeks, or for months to years. Older people coming to the end of their life without illness may have some of the same care issues. They can also benefit from the approaches to care taken in palliative care.

Common care issues in palliative care include:

- pain
- dyspnoea (breathing difficulty)
- dysphagia (difficulty swallowing)
- constipation/incontinence (bowel and/or bladder management)

Who should attend a case conference?

Staff in residential aged care facilities and providers of home care often meet with families. If possible, the person receiving care should attend, their GP, and any concerned family members or friends.

Your contact for this case conference is:

Name of staff member:

Telephone:

- anxiety

Organisation:

• fatigue (tiredness).

Invitation for you and your family

Organisation:

Palliative care case conference

A palliative care case conference has been organised for:

Name of resident/client:	
Resident/client date of birth (dd/mm/yy):	
Case conference date (dd/mm/yy):	Start time:
Location:	

Please let us know if you can attend. If you would like to join by telephone, let us know and provide a suitable number to contact you.

Your contact for this case conference is:

Name of staff member:		
Role:		
Telephone:		



On the next page you will find information on palliative care and palliative care case conferences

Invitation for you and your family

Palliative care case conference

Case conference: Case conferences or family meetings are an opportunity to discuss a person's care needs. They ideally include the person (if able to attend), their family and/or their substitute decision-maker, and members of the care team including the doctor.

Palliative care: Palliative care is person- and family-centred care that supports a person to live the best life they can with a life-limiting illness. This means that the person has little or no prospect of cure and is expected to die. The focus is on quality of life.

Life-limiting illnesses include dementia, advanced heart, kidney, lung or liver disease, cancer, and motor neurone disease.

People can receive palliative care for days or weeks, or for months to years. Older people coming to the end of their life without illness may have some of the same care issues. They can also benefit from the approaches to care taken in palliative care.

Common care issues in palliative care include:

- pain
- dyspnoea (breathing difficulty)
- dysphagia (difficulty swallowing)
- constipation/incontinence (bowel and/or bladder management)

- depression
- delirium (sudden confusion)
- anxiety
- nausea (feel that you want to vomit)
- fatigue (tiredness).

Who should attend a case conference?

Staff in residential aged care facilities and providers of home care often meet with families. If possible, the person receiving care should attend, their GP, and any concerned family members or friends.

Confirmation	for	you	and
your family			

Organisation:

					<i>c</i>
Ρ	้อเ	liative	care	case	conference
	-			0000	

A palliative care case conference has been organised for:

Name of resident/client:	
Resident/client date of birth (dd/mm/yy):	
Case conference date (dd/mm/yy):	Start time:
Location:	

Your involvement in planning care is important. If you are unable to attend in person but would like to join by telephone, please dial in using the following telephone number and code.

Dial-in telephone number:	Code:
Your contact for this case conference is:	
Name of staff member:	
Role:	
Telephone:	

Please write down if there are any issues you want to talk about and remember to bring this form with you to the meeting so that this can be included.

Staff communication sheet

Palliative care case conference

A palliative care case conference has been organised for:

Name of resident/client:	
Case conference date (dd/mm/yy):	Start time:
Location:	

As valuable members of the care team your contribution to the case conference is important. Please list below any issues, concerns or suggestions you would like mentioned. Common issues include review of symptoms (e.g. pain, dyspnoea), concerns with nutrition or hydration, family issues, emotional concerns of the resident. If you are available and would like to attend the case conference, please contact the Case Conference Facilitator:

Name of facilitator:

lssue, concern or suggestion. Please be as specific as possible.	Designation

Organisation:

Palliative care case conference

Full name of client:	
DOB (dd/mm/yy):	
Purpose of case conference:	

Client consent/substitute decision-maker (SDM) consent

.

My care provider has explained the purpose of a case conference and I give permission for my care provider to prepare a case conference. I give permission to the providers listed below to participate in the case conference and discuss my/my family member's medical history, diagnosis, and current needs.

Signature:		
Date:		

Dial-in telephone number:	Ĺ	.ode:
Client in attendance? Yes	No If no, give reason:	
Family members		
Name	Relationship	Attending in person (P) or teleconference (T)
		РТ
Health and care professionals		
Name	Discipline/position	Attending in person (P) or teleconference (T)
		РТ

D¹ **I I I I**

Palliative care case conference

Start time:

Need (as appropriate):

Key Issues	Description
Advance care plan	
Does this need to be reviewed? Does the person understand their diagnosis/prognosis?	
Symptoms	
For example: fatigue, anorexia, pain, nausea, dyspnoea, dysphagia	
Social/psychological needs	
For example: isolation, anxiety, depression What supports are being provided? What supports are needed?	
Assessments/investigations	
Can the client manage ADL's (Activities of Daily Living)? Do they need additional support?	
Carer/family issues or needs	
For example: has a Needs Assessment Tool for Carers (NAT-C) been completed?	
Other	
For example: general issues, housing issues, financial issues	

Palliative care case conference

Agreed action plan

Goal	Actions	Key person(s) responsible	Description

Palliative care case conference

Time	completed:
Gene	ral practitioner:
Tick a	ppropriate box
	Original placed in the client's clinical notes
	Copy provided to all participants
	Copy sent to GP
	Client's care plan and assessment reviewed and updated

Palliative care case conference facilitator

Name:	Position:	
Signature:	Date (dd/mm/yy):	



PBS Prescriber Bag medicines for terminal phase symptoms

These medicines are available through the PBS at no cost to prescribers. They can be provided free to patients during home visits for emergency use in managing symptoms or bridging the gap until a prescription is dispensed. Medicines shaded are part of the National Core Community Palliative Care Medicines List and are used to treat common symptoms in straightforward cases.

Medicine	Clinical uses in terminal phase	Strength and form	Pack size	Max qty (packs)
Adrenaline (Epinephrine)	Airway obstruction (nebulised), small volume malignant bleeding (topical)	1 in 1000 (1 mg/mL) injection	5 x 1mL	1
Clonazepam	Agitation, anxiety, distressing breathlessness, refractory distress, seizure	2.5 mg/mL drops	1 x 10mL	1
Furosemide	Oedema associated with heart failure	20 mg/2 mL injection	5 x 2mL	1
Haloperidol	Anxiety, delirium, nausea/vomiting, refractory distress, terminal restlessness			1
Hydrocortisone	Acute severe breathlessness/spinal cord compression, in place of dexamethasone	100 or 250 mg injection (reconstituted to 2mL)	1 x dual chamber vial	2 (100mg) or 1 (250mg)
Hyoscine butylbromide	Respiratory tract secretions, noisy breathing, managing cramps with bowel obstruction	20 mg/mL injection	5 x 1mL	1
Metoclopramide	Nausea/vomiting	10 mg/2 mL injection	10 x 2mL	1
Midazolam	Agitation, distressing breathlessness, refractory distress, seizure	5 mg/mL injection	10 x 1mL	1
Morphine	Distressing breathlessness (first line), pain	10, 15, 20, or 30 mg/mL injection	5 x 1mL	1
Naloxone	Reversing life-threatening opioid overdose	400 microgram/mL injection	5 or 10 x 1mL	2

Based on: caring@home/Pharmaceutical Society of Australia. National Core Community Palliative Care Medicines List [Internet]. Brisbane, QLD: caring@home; 2024 [cited 2025 Jun 2]. Available from: <u>https://www.caringathomeproject.com.au/for-health-professionals/national-core-community-palliative-care-medicines-list</u>



The PBS Prescriber Bag for palliative care

People with palliative care needs may choose to be cared for and die at home. This may include their private dwelling or a residential aged care facility. The evidence encourages the prescribing of all terminal phase medicines in advance, known as anticipatory prescribing. While it should not be a substitute for good advance planning, the PBS Prescriber Bag provides a safety net for those who deteriorate rapidly and unexpectedly at the end of life. This ensures rapid symptom management when needed, though deterioration can occur suddenly.

Prescriber Bag supply order forms allow monthly ordering of medicines and can be requested from Services Australia: <u>https://www.servicesaustralia.gov.au/pbs-and-rpbs-official-stationery?</u> <u>context=20</u>.

Complete the form, sign it, and give it to a community pharmacist for dispensing.

Dosing information

For specific dosing advice, refer to the CareSearchgp or palliMEDS apps (free to download), Palliative Care Therapeutic Guidelines, or Australian Medicines Handbook. You can also consult your local pharmacist.

Notes on use of specific medicines

- Morphine: avoid repeated dosing in people with serious kidney failure.
- Clonazepam or midazolam: may help with breathlessness if anxiety is present. They may also help to relieve rigidity associated with end-stage Parkinson's Disease if dopaminergic medication has ceased.
- Adrenaline in nebulised form may give temporary relief of stridor with breathlessness.

Practical tips

- Order your PBS Prescriber Bag medicines at the end of each month.
- Securely store S8 medicines (especially opioids) and follow local legislative guidelines.
- Consider carrying equipment to administer medicines subcutaneously.
- Limit subcutaneous injections to 1.5 mL to avoid causing pain at the injection site.
- Keep a notepad to record medication administration and any doses discarded.

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Supporting a planned home death: The GP's Checklist

The GP has a critical role in end of life care for patients who wish to die at home. This checklist guides GPs through decision-making. care of the patient and family, and identifying the need for appropriate supports early so that issues can be addressed ahead of time. GPs managing patients dying at home usually share care with other services, including palliative care and home nursing. This checklist can act as a planning tool for shared care, and a trigger to help clarify how care will be organised between those involved.

atie	ent name/ID: Date:
	1 Clarify expectations and support
	Has the patient indicated they want to die at home?
	Actions needed:
	Do those who live with the patient know about and share that wish?
	 Has the plan been discussed within the family? Consider – young children, others with care needs in the household
	Actions needed:
	Are there enough people to share the care?
	• Consider practical, hands-on availability for round the clock care. Suggest a roster to support carer and provide time out.
	• Consider specific services that can support families caring for someone who is dying at home, e.g.
	night nursing services or volunteers. The local palliative care service can advise.
	Actions needed:

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Review the patient's prognosis, in order to help the family plan ahead. E.g. Is care likely to be needed for days, weeks, or months?

Actions needed:

Is there a back-up plan if either the patient or the family find it difficult?

- Clarify and document a plan and ensure that it is realistic, and understood by all involved.
- Where appropriate, provide a letter (or Ambulance Plan) describing the palliative goals of care in case of a triple zero call. Clearly state that the patient is dying and cardiopulmonary resuscitation is not appropriate, where agreed.

Actions needed:

2 Assess the home situation

Can the patient be cared for safely and comfortably in the home?

- Refer to home nursing services, and ask them to teach the family about how to provide care safely (transferring and moving the person, eating and drinking, giving medicines). How much nursing support is available? Specifically, how many visits can the patient have?
- Are there complex nursing needs that will be difficult to manage at home e.g. difficult wounds, fistulas, spinal analgesia?
- Consider equipment for nursing a bed-bound patient. They will need a hospital bed, mobility aids, commodes and personal care equipment, wheelchair, pressure mattresses, etc. Consider a palliative care referral for OT and/or physiotherapy assessment to advise on and organise equipment.
- Discuss the option of an in-dwelling catheter to reduce the care burden for a bedbound patient.
- Encourage the family to think about practical arrangements to make caring easier or safer e.g. moving a patient's bed to a different room, or patient moving in with a family member (remember though that moving to a different address can disrupt eligibility for services – so do this early if possible.)

Actions needed:

3 Plan for symptom management

Review long-term medications. Cease any that no longer contribute to patient's comfort.

Actions	needed:
Actions	necucu.

Discuss with the family how the patient's symptoms will be reviewed and managed.

- How often GP visits will occur
- What the home nurses will do
- What the palliative care service will do
- Arrangements to provide prescriptions

Actions needed:

Plan for predictable, common symptoms that occur at the end of life:

- Dyspnoea / terminal secretions
- Delirium

- Pain
- Nausea

Actions needed:

Ensure emergency medications are available in the house for when they are needed. This is best done well in advance as deterioration can be unpredictable.

- Remember dying patients cannot take oral medications
- Subcutaneous medications are preferred to ensure continuing symptom control, with bolus medications via sc butterfly needle, and/or a syringe driver with a 24 hour infusion
- Family members should be taught how to give breakthrough doses by palliative or home nurses
- Check that medications are available at a community pharmacy, and that the caregivers have an adequate supply to get through after hours and weekends in particular

Actions needed:

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- Consider whether a plan is needed for high risk problems such as major bleed, airway or bowel obstruction.
- If care needs are complex, or a high risk problem exists, seek early advice from a palliative care specialist.

Actions needed:

4 Information that families need

Are the carers fully prepared for the fact that the dying person will be dependent and bedbound? Actions needed:

Do the carers need information about eating and drinking in the palliative care situation? Do they know that loss of appetite is a common and predictable feature of advanced disease? Are they aware that wallowing deteriorates with the approach of the terminal phase? Do they need ideas about what and how much to offer the patient to eat and drink, and how to do this safely?

Actions needed:

Do the carers need information about physical changes that occur as a person is dying, including:

- Changes in breathing patterns, including the possibility of terminal secretions ('death rattle')
- Changes in skin colour and temperature
- Changes in level of consciousness, including the possibility of terminal delirium

Actions needed:

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Ensure that the family has access to 24-hour phone advice about symptoms or changes in the patient's condition. Make sure these contact numbers are accessible to anyone who might need them. Actions needed:

Do the caregivers need information about what to do after the patient dies?

- Encourage them to think about choosing a funeral director
- Reassure them that there is no urgency to ring anyone straight away after the patient dies
- Ensure that they know which doctor has agreed to certify death, and the arrangements for contacting them.

Actions needed:

Facility name:			
Address:			

For more information visit CareSearch GP Hub www.caresearch.com.au Here you will find family resources you can order and give to patients and carers.



MBS items supporting nurse practitioner palliative care provision

MBS Item	Activities	MBS Benefit 100% (as of 01/08/24)
<u>82200</u>	Brief attendance	\$14.20
	Professional attendance by a participating nurse practitioner for an obvious problem characterised by the straightforward nature of the task that requires a short patient history and, if required, limited examination and management.	
	Telehealth: Item <u>91192</u> . Phone: Item <u>91193</u> .	
<u>82205</u>	Short attendance (<20 mins)	\$31.05
	Includes any of the following: taking a history; undertaking clinical examination; arranging any necessary investigation; implementing a management plan; or providing appropriate preventive health care, for one or more health related issues, with appropriate documentation.	
	Telehealth: Item <u>91178</u> . Phone: Item <u>91189</u> .	
<u>82210</u>	Detailed consultation (>20 mins)	\$58.85
	Includes any of the following: taking a detailed history; undertaking clinical examination; arranging any necessary investigation; implementing a management plan; or providing appropriate preventive health care, for one or more health related issues, with appropriate documentation.	
	Telehealth: Item <u>91179</u> . Phone: Item <u>91190</u> .	
<u>82215</u>	Extensive consultation (>40 mins)	\$86.80
	Includes any of the following: taking an extensive history; undertaking clinical examination; arranging any necessary investigation; implementing a management plan; or providing appropriate preventive health care, for one or more health related issues, with appropriate documentation.	
	Telehealth: Item <u>91180</u> . Phone: Item <u>91191</u> .	



MBS items supporting a planned general practice palliative care pathway

This document uses the MBS items currently available to general practitioners to suggest a timeframe and care pathway for community patients with palliative care needs. A separate pathway is provided for residential aged care patients (398kb pdf)

Suggested timeframe	Medicare initiative	Activities	MBS Item		MBS Benefit 100% (as of 01/08/24)
0 months	Over 75 years health assessment	An annual assessment of a patient's health and physical,	<u>701</u> (<30 mins)		\$67.60
		psychological, and social function. Select relevant item based on complexity and Practice Nurse and GP time. An opportunity to initiate	<mark>703</mark> (30-45 mins)		\$157.10
			<mark>705</mark> (45-60 mins)		\$216.80
		discussion of advance care planning or palliative care.		<u>)7</u> mins)	\$306.25
2nd week	GP management plan (GPMP)	For managing terminal medical conditions by preparing, coordinating, reviewing, or contributing to chronic disease management plans. An opportunity to introduce advance care planning or to discuss palliative care.	<u>721</u> <u>723</u>		\$164.35
	Team care arrangement (TCA) coordination	Requires at least two collaborating providers, each providing a different kind of treatment (e.g., allied health services or medical practitioner).			\$130.25
3rd week	GP Mental Health Treatment Plan	For preparing a mental health care plan for a patient with mental health care needs.	No GP mental health	2700 (20-40 mins)	\$81.70
		Select relevant item depending on time and GP training.	training	<u>2701</u> (>40 mins)	\$120.25

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Suggested Timeframe	Medicare initiative	Activities	MBS Item		MBS Benefit 100% (as of 01/08/24)	
		As per the Diagnostic and Statistical Manual of Mental Disorders (DSM-V-TR)	GP mental health training	2715 (20-40 mins)	\$103.70	
				<u>2717</u> (>40 mins)	\$152.80	
1st month	GP-organised and coordinated multidisciplinary care conference	An opportunity for an holistic informed approach to ongoing care. Covers organising, coordinating, and participating in a case conference involving the patient, family/significant others, and a minimum of two other health providers.	<u>739</u> (20-40 mins)		\$137.75	
2nd month	Domiciliary Medication Management Review (DMMR)	Referral to eligible pharmacist. Ensures optimal management of patients with multiple medications and/or complexity.	<u>900</u>		\$176.40	
4th month	Level D or E consultation	An extended consultation for the purpose of managing palliative care and end-of-life	44 (Level D: >40 mins) <u>123</u> (Level E: >60 mins)		\$122.15	
		care needs, discussing goals of care, advance care planning, or for completing an Advance Care Directive based on earlier discussions.			\$197.90	
5th month	Review GP Mental Health Plan	4 weeks to 6 months after preparation of plan, review referral feedback and progress against goals.	<u>2712</u>		\$81.70	
6th month	Review of GPMP or TCA	Discuss progress against goals and actions.	732		\$82.10	
8th month	GP-led case conference	A real-time discussion of any changing needs and goals of care with patient, significant others, and the health care team.	<u>735</u> (15-20 mins)		\$80.55	
5 per year	Practice nurse care plan monitoring	Where a GPMP or TCA is in place.	<u>10997</u>		\$13.65	

Based on: PHN North Western Melbourne. MBS remuneration to support planned palliative care for patients: A guide for health professionals working in general practice and residential aged care. Melbourne: NWMPHN; 2017 [cited 2024 May 27]. Available from: <u>https://nwmphn.org.au/wp-content/uploads/2020/12/NWMPHN-</u> Palliative-Care-For-GP-and-RAC5.pdf