

## 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.

atien	tient 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?	
,	<ul> <li>Has the plan been discussed within the family?</li> </ul>	
,	<ul> <li>Consider – young children, others with care needs in the household</li> </ul>	
	Actions needed:	
	Are there enough people to share the care?	
•	<ul> <li>Consider practical, hands-on availability for round the clock care. Sugge and provide time out.</li> </ul>	est a roster to support carei
•	<ul> <li>Consider specific services that can support families caring for someone night nursing services or volunteers. The local palliative care service can</li> </ul>	
,	Actions needed:	

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Ac	tions needed:
ls t	here 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 car
	case of a triple zero call. Clearly state that the patient is dying and cardiopulmonary
	resuscitation is not appropriate, where agreed.
Ac	tions needed:
	Assess the home situation  In the patient be cared for safely and comfortably in the home?  Refer to home pursing services, and ask them to took the family about how to provide sare.
•	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 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 patien
•	Encourage the family to think about practical arrangements to make caring easier or safer – e 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 d this early if possible.)
۸۰۰	tions needed:
	alons necucu.

Discuss with the family how the patient's sympto	ms will be reviewed and managed.
How often GP visits will occur	
<ul> <li>What the home nurses will do</li> </ul>	
What the palliative care service will do	
Arrangements to provide prescriptions	
Actions needed:	
Plan for predictable, common symptoms that occ	
Dyspnoea / terminal secretions	• Pain
• Delirium	Nausea
Actions needed:	
Ensure emergency medications are available in th	ne house for when they are needed. This is hes
well in advance as deterioration can be unpredict	·
<ul> <li>Remember – dying patients cannot take oral i</li> </ul>	
. •.	nsure continuing symptom control, with bolus
medications via sc butterfly needle, and/or a	
•	e breakthrough doses by palliative or home nu
	nmunity pharmacy, and that the caregivers have
adequate supply to get through after hours a	
	nmunity pharmacy, and that the caregivers ha

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obstruction.	ether a plan is needed for high risk problems such as major bleed, airway or bowel
	eds are complex, or a high risk problem exists, seek early advice from a palliative car
Actions need	ed:
4 Inforr	nation that families need
Are the carers	s fully prepared for the fact that the dying person will be dependent and bedbound?
	need information about eating and drinking in the palliative care situation? Do the ppetite is a common and predictable feature of advanced disease? Are they aware to
that loss of a wallowing de	ppetite is a common and predictable feature of advanced disease? Are they aware teriorates with the approach of the terminal phase? Do they need ideas about what offer the patient to eat and drink, and how to do this safely?
that loss of a wallowing de how much to Actions need	ppetite is a common and predictable feature of advanced disease? Are they aware teriorates with the approach of the terminal phase? Do they need ideas about what offer the patient to eat and drink, and how to do this safely?
that loss of a wallowing de how much to  Actions need  Do the carers  Changes i	ppetite is a common and predictable feature of advanced disease? Are they aware teriorates with the approach of the terminal phase? Do they need ideas about what offer the patient to eat and drink, and how to do this safely?  ed:  need information about physical changes that occur as a person is dying, including:

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	ure that the family has access to 24-hour phone advice about symptoms or changes in the ient's condition. Make sure these contact numbers are accessible to anyone who might need
Acti	ons needed:
	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.
Act	ions needed:
Fac	ility name:
Δ٨٥	dress:

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.