

Essential Component 5: Care is based on the assessed needs of the patient, carer and family – standards comparison

National Palliative Care Standards, 5th Edition

Relevant standard	Relevant action (where available) against Essential Component 5
<p>1. Assessment of needs Initial and ongoing assessment incorporates the person's physical, psychological, cultural, social and spiritual experiences and needs</p>	<p>1.1 The initial and ongoing assessments are carried out by qualified interdisciplinary personnel.</p> <p>1.3 Clinical assessment tools are informed by the best available evidence and identify those approaching the end-of-life as well as those that are imminently dying.</p> <p>1.4 The person's needs are reassessed on a regular basis.</p> <p>1.6 Ongoing assessments are used to inform the care plan and any subsequent changes to it.</p>
<p>2. Developing the care plan The person, their family and carers work in partnership with the team to communicate, plan, set goals of care and support informed decisions about the care plan</p>	<p>2.1 Care planning is informed by the assessment process and reflects a person-centred, holistic approach that incorporates cultural, spiritual, physical, psychological and social needs.</p> <p>2.6 Specific attention is paid to the needs of people who may be vulnerable or at risk, to support communication, goal setting and care planning. This includes but is not limited to Aboriginal and Torres Strait Islanders, asylum seekers, people who have experienced torture and trauma, people who are experiencing homelessness, people living with mental illness, intellectual disabilities or dementia, paediatric populations or people from ethnically and culturally diverse backgrounds.</p> <p>2.8 The expectations and preferences of the person, their family and carers for the type and place of care are discussed, negotiated and an agreed plan is documented.</p> <p>2.9 The care plan is reviewed and updated regularly, based on reassessments of the person's condition, needs, and preferences, and in consultation with the person, their family and carers. Changes to the care plan are documented.</p> <p>2.10 Care plans incorporate management for emergency and after-hours support, including certification of death and plans for the care and collection of the body where this is required after hours.</p>

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<p>3. Caring for carers</p> <p>The needs and preferences of the person's family and carers are assessed, and directly inform provision of appropriate support and guidance about their role</p>	<p>3.1 At least one carer is identified for each person as far as possible and their specific needs, including their need for information, are assessed and documented.</p> <p>3.2 The service works with the family and carers to understand their needs and desired level of involvement in care. The potential benefits and risks around assisting with care are discussed with the person, their family and carers and there is ongoing assessment of their willingness and ability to participate in the provision of care.</p> <p>3.4 The family and carers are provided with up-to-date information and resources that are adapted to meet their needs and that inform their participation in care planning and delivery. This may include information about accessing respite services, equipment, financial support and other services, as well as encouraging the involvement of personal support networks and self-care.</p>
<p>4. Providing care</p> <p>The provision of care is based on the assessed needs of the person, informed by evidence, and is consistent with the values, goals and preferences of the person as documented in their care plan</p>	<p>4.1 Care is delivered promptly, in accordance with the changing needs of the person, their family and carers, their documented care plan and their goals and preferences.</p> <p>4.3 Where care cannot be delivered in accordance with the goals and preferences of the person, this is discussed with the person, their family, and carers, and an agreed alternative plan is documented and communicated.</p> <p>4.4 There are protocols and procedures in place for the escalation of care where required, based on assessed needs.</p>

National Safety and Quality Health Service (NSQHS) Standards, 2nd Edition

Relevant standard	Relevant action (where available) against Essential Component 5
<p>1. Clinical governance</p> <ul style="list-style-type: none"> • Patient safety and quality systems • Clinical performance and effectiveness 	<p>1.15 The health service organisation:</p> <ul style="list-style-type: none"> a. identifies the diversity of the consumers using its services b. identifies groups of patients using its services who are at higher risk of harm c. incorporates information on the diversity of its consumers and higher risk groups into the planning and delivery of care. <p>1.20 The health service organisation uses its training systems to:</p> <ul style="list-style-type: none"> a. assess the competency and training needs of its workforce b. implement a mandatory training program to meet its requirements arising from these standards c. provide access to training to meet its safety and quality training needs d. monitor the workforce's participation in training.

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	<p>1.21 The health service organisation has strategies to improve the cultural awareness and cultural competency of the workforce to meet the needs of its Aboriginal and Torres Strait Islander patients.</p> <p>1.23 The health service organisation has processes to:</p> <ol style="list-style-type: none"> a. define the scope of clinical practice for clinicians, considering the clinical service capacity of the organisation and clinical services plan b. monitor clinicians’ practices to ensure that they are operating within their designated scope of clinical practice c. review the scope of clinical practice of clinicians periodically and whenever a new clinical service, procedure or technology is introduced or substantially altered. <p>1.24 The health service organisation:</p> <ol style="list-style-type: none"> a. conducts processes to ensure that clinicians are credentialed, where relevant b. monitors and improves the effectiveness of the credentialing process. <p>1.26 The health service organisation provides supervision for clinicians to ensure that they can safely fulfil their designated roles, including access to after-hours advice, where appropriate.</p> <p>1.27 The health service organisation has processes that:</p> <ol style="list-style-type: none"> a. provide clinicians with ready access to best-practice guidelines, integrated care pathways, clinical pathways and decision support tools relevant to their clinical practice b. support clinicians to use the best available evidence, including relevant clinical care standards developed by the Australian Commission on Safety and Quality in Health Care.
<p>2. Partnering with consumers</p> <ul style="list-style-type: none"> • Partnering with patients in their own care • Health literacy 	<p>2.6 The health service organisation has processes for clinicians to partner with patients or their substitute decision-maker to plan, communicate, set goals, and make decisions about their current and future care.</p> <p>2.7 The health service organisation supports the workforce to form partnerships with patients and carers so that patients can be actively involved in their own care.</p> <p>2.8 The health service organisation uses communication mechanisms that are tailored to the diversity of the consumers who use its services and, where relevant, the diversity of the local community.</p>

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<ul style="list-style-type: none"> Partnering with consumers in organisational design and governance 	<p>2.10 The health service organisation supports clinicians to communicate with patients, carers, families and consumers about health and health care so that:</p> <ol style="list-style-type: none"> information is provided in a way that meets the needs of patients, carers, families and consumers information provided is easy to understand and use the clinical needs of patients are addressed while they are in the health service organisation information needs for ongoing care are provided on discharge. <p>2.13 The health service organisation works in partnership with Aboriginal and Torres Strait Islander communities to meet their healthcare needs.</p>
<p>3. Preventing and controlling healthcare-associated infection</p> <ul style="list-style-type: none"> Infection prevention and control systems 	<p>3.6 Clinicians assess infection risks and use transmission-based precautions based on the risk of transmission of infectious agents, and consider:</p> <ol style="list-style-type: none"> patients' risks, which are evaluated at referral, on admission or on presentation for care, and re-evaluated when clinically required during care whether a patient has a communicable disease, or an existing or a pre-existing colonisation or infection with organisms of local or national significance accommodation needs to manage infection risks the need to control the environment precautions required when the patient is moved within the facility or to external services the need for additional environmental cleaning or disinfection equipment requirements.
<p>4. Medication safety</p> <ul style="list-style-type: none"> Documentation of patient information Continuity of medication management 	<p>4.5 Clinicians take a best possible medication history, which is documented in the healthcare record on presentation or as early as possible in the episode of care.</p> <p>4.6 Clinicians review a patient's current medication orders against their best possible medication history and the documented treatment plan and reconcile any discrepancies on presentation and at transitions of care.</p> <p>4.10 The health service organisation has processes:</p> <ol style="list-style-type: none"> to perform medication reviews for patients, in line with evidence and best practice to prioritise medication reviews, based on a patient's clinical needs and minimising the risk of medication-related problems that specify the requirements for documentation of medication reviews, including actions taken as a result.

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<ul style="list-style-type: none"> Delivering comprehensive care 	<p>5.13 Clinicians use processes for shared decision making to develop and document a comprehensive and individualised plan that:</p> <ol style="list-style-type: none"> addresses the significance and complexity of the patient’s health issues and risks of harm identifies agreed goals and actions for the patient’s treatment and care identifies the support people a patient wants involved in communications and decision-making about their care commences discharge planning at the beginning of the episode of care includes a plan for referral to follow-up services, if appropriate and available is consistent with best practice and evidence. <p>5.14 The workforce, patients, carers and families work in partnership to:</p> <ol style="list-style-type: none"> use the comprehensive care plan to deliver care monitor the effectiveness of the comprehensive care plan in meeting the goals of care review and update the comprehensive care plan if it is not effective reassess the patient’s needs if changes in diagnosis, behaviour, cognition, or mental or physical condition occur. <p>5.15 The health service organisation has processes to identify patients who are at the end-of-life that are consistent with the <i>National Consensus Statement: Essential elements for safe and high-quality end-of-life care</i>.</p> <p>5.20 Clinicians support patients, carers and families to make shared decisions about end-of-life care in accordance with the <i>National Consensus Statement: Essential elements for safe and high-quality end-of-life care</i>.</p>
<ul style="list-style-type: none"> Minimising patient harm 	<p>5.21 The health service organisation providing services to patients at risk of pressure injuries has systems for pressure injury prevention and wound management that are consistent with best-practice guidelines.</p> <p>5.22 Clinicians providing care to patients at risk of developing, or with, a pressure injury conduct comprehensive skin inspections in accordance with best-practice time frames and frequency.</p> <p>5.23 The health service organisation providing services to patients at risk of pressure injuries ensures that:</p> <ol style="list-style-type: none"> patients, carers and families are provided with information about preventing pressure injuries

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	<p data-bbox="571 286 1444 398">b. equipment, devices and products are used in line with best-practice guidelines to prevent and effectively manage pressure injuries.</p> <p data-bbox="523 443 1428 555">5.24 The health service organisation providing services to patients at risk of falls has systems that are consistent with best-practice guidelines for:</p> <ul style="list-style-type: none"> <li data-bbox="571 555 821 589">a. falls prevention <li data-bbox="571 593 965 627">b. minimising harm from falls <li data-bbox="571 631 917 665">c. post-fall management. <p data-bbox="523 710 1436 822">5.25 The health service organisation providing services to patients at risk of falls ensures that equipment, devices and tools are available to promote safe mobility and manage the risks of falls.</p> <p data-bbox="523 866 1460 978">5.26 Clinicians providing care to patients at risk of falls provide patients, carers and families with information about reducing falls risks and falls prevention strategies.</p> <p data-bbox="523 1023 1452 1169">5.27 The health service organisation that admits patients overnight has systems for the preparation and distribution of food and fluids that include nutrition care plans based on current evidence and best practice.</p> <p data-bbox="523 1214 1460 1281">5.28 The workforce uses the systems for preparation and distribution of food and fluids to:</p> <ul style="list-style-type: none"> <li data-bbox="571 1292 1268 1326">a. meet patients’ nutritional needs and requirements <li data-bbox="571 1330 1204 1364">b. monitor the nutritional care of patients at risk <li data-bbox="571 1368 1436 1435">c. identify, and provide access to, nutritional support for patients who cannot meet their nutritional requirements with food alone <li data-bbox="571 1440 1348 1507">d. support patients who require assistance with eating and drinking. <p data-bbox="523 1563 1452 1675">5.29 The health service organisation providing services to patients who have cognitive impairment or are at risk of developing delirium has a system for caring for patients with cognitive impairment to:</p> <ul style="list-style-type: none"> <li data-bbox="571 1686 1452 1832">a. incorporate best-practice strategies for early recognition, prevention, treatment and management of cognitive impairment in the care plan, including the <i>delirium clinical care standard</i>, where relevant <li data-bbox="571 1836 1396 1904">b. manage the use of antipsychotics and other psychoactive medicines, in accordance with best practice and legislation. <p data-bbox="523 1948 1388 2060">5.30 Clinicians providing care to patients who have cognitive impairment or are at risk of developing delirium use the system for caring for patients with cognitive impairment to:</p> <ul style="list-style-type: none"> <li data-bbox="571 2072 1396 2094">a. recognise, prevent, treat and manage cognitive impairment

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	<p>b. collaborate with patients, carers and families to understand the patient and implement individualised strategies that minimise any anxiety or distress while they are receiving care.</p> <p>5.31 The health service organisation has systems to support collaboration with patients, carers and families to:</p> <ol style="list-style-type: none"> a. identify when a patient is at risk of self-harm b. identify when a patient is at risk of suicide c. safely and effectively respond to patients who are distressed, have thoughts of self-harm or suicide, or have self-harmed. <p>5.33 The health service organisation has processes to identify and mitigate situations that may precipitate aggression.</p> <p>5.34 The health service organisation has processes to support collaboration with patients, carers and families to:</p> <ol style="list-style-type: none"> a. identify patients at risk of becoming aggressive or violent b. implement de-escalation strategies c. safely manage aggression, and minimise harm to patients, carers, families and the workforce. <p>5.35 Where restraint is clinically necessary to prevent harm, the health service organisation has systems that:</p> <ol style="list-style-type: none"> a. minimise and, where possible, eliminate the use of restraint b. govern the use of restraint in accordance with legislation c. report use of restraint to the governing body. <p>5.36 Where seclusion is clinically necessary to prevent harm and is permitted under legislation, the health service organisation has systems that:</p> <ol style="list-style-type: none"> a. minimise and, where possible, eliminate the use of seclusion b. govern the use of seclusion in accordance with legislation c. report use of seclusion to the governing body.
<p>6.Communicating for safety</p> <ul style="list-style-type: none"> • Clinical governance and quality improvement to support effective communication • Communication at clinical handover 	<p>6.3 Clinicians use organisational processes from the <i>Partnering with Consumers Standard</i> to effectively communicate with patients, carers and families during high-risk situations to:</p> <ol style="list-style-type: none"> a. actively involve patients in their own care b. meet the patient’s information needs c. share decision-making. <p>6.7 The health service organisation, in collaboration with clinicians, defines the:</p> <ol style="list-style-type: none"> a. minimum information content to be communicated at clinical handover, based on best-practice guidelines

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<ul style="list-style-type: none"> • Communication of critical information • Documentation of information 	<ul style="list-style-type: none"> b. risks relevant to the service context and the particular needs of patients, carers and families c. clinicians who are involved in the clinical handover. <p>6.9 Clinicians and multidisciplinary teams use clinical communication processes to effectively communicate critical information, alerts and risks, in a timely way, when they emerge or change to:</p> <ul style="list-style-type: none"> a. clinicians who can make decisions about care b. patients, carers and families, in accordance with the wishes of the patient. <p>6.10 The health service organisation ensures that there are communication processes for patients, carers and families to directly communicate critical information and risks about care to clinicians.</p> <p>6.11 The health service organisation has processes to contemporaneously document information in the healthcare record, including:</p> <ul style="list-style-type: none"> a. critical information, alerts and risks b. reassessment processes and outcomes c. changes to the care plan.
<p>8. Recognising and responding to acute deterioration</p> <ul style="list-style-type: none"> • Detecting and recognising acute deterioration and escalating care 	<p>8.4 The health service organisation has processes for clinicians to detect acute physiological deterioration that require clinicians to:</p> <ul style="list-style-type: none"> a. document individualised vital sign monitoring plans b. monitor patients as required by their individualised monitoring plan c. graphically document and track changes in agreed observations to detect acute deterioration over time, as appropriate for the patient. <p>8.5 The health service organisation has processes for clinicians to recognise acute deterioration in mental state that require clinicians to:</p> <ul style="list-style-type: none"> a. monitor patients at risk of acute deterioration in mental state, including patients at risk of developing delirium b. include the person’s known early warning signs of deterioration in mental state in their individualised monitoring plan c. assess possible causes of acute deterioration in mental state, including delirium, when changes in behaviour, cognitive function, perception, physical function or emotional state are observed or reported d. determine the required level of observation e. document and communicate observed or reported changes in mental state. <p>8.6 The health service organisation has protocols that specify criteria for escalating care, including:</p>

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<ul style="list-style-type: none"> Responding to acute deterioration 	<ul style="list-style-type: none"> a. agreed vital sign parameters and other indicators of physiological deterioration b. agreed indicators of deterioration in mental state c. agreed parameters and other indicators for calling emergency assistance d. patient pain or distress that is not able to be managed using available treatment e. worry or concern in members of the workforce, patients, carers and families about acute deterioration. <p>8.7 The health service organisation has processes for patients, carers or families to directly escalate care.</p> <p>8.8 The health service organisation provides the workforce with mechanisms to escalate care and call for emergency assistance.</p> <p>8.9 The workforce uses the recognition and response systems to escalate care.</p> <p>8.10 The health service organisation has processes that support timely response by clinicians with the skills required to manage episodes of acute deterioration.</p> <p>8.11 The health service organisation has processes to ensure rapid access at all times to at least one clinician, either on site or in close proximity, who can deliver advanced life support.</p> <p>8.12 The health service organisation has processes to ensure rapid referral to mental health services to meet the needs of patients whose mental state has acutely deteriorated.</p> <p>8.13 The health service organisation has processes for rapid referral to services that can provide definitive management of acute physical deterioration.</p>

National Consensus Statement (Adult and Paediatric)

Relevant standard	Relevant action (where available) against Essential Component 5
<p>1. Patient/family centred care Patients/children and families are part of decision making about end-of-life care</p>	<p>Adult</p> <p>1.2 The clinical team should work with the patient, family and carers to identify the substitute decision-maker, family spokesperson or other key contacts that the patient wishes to be involved in discussions about their care.</p>

Relevant standard	Relevant action (where available) against Essential Component 5
	<p>1.3 In some cultures, mainstream assumptions about death and dying, and about patients and families as decision-makers may not be correct. Culturally appropriate decision makers should be identified as early as possible so that strategies can be put in place for obtaining their input into discussions about end-of-life care.</p> <p>1.4 Clinicians should seek to understand, and be respectful, sensitive and responsive to, the individual preferences and needs of all patients, substitute decision-makers, families and carers, regardless of aspects of identity such as culture, religious belief, gender or sexual preference.</p> <p>1.5 Whenever possible, clinicians should prepare for having conversations about end-of-life care. Necessary preparation may include:</p> <ul style="list-style-type: none"> • reaching consensus among all of the clinical teams involved in the patient’s care about the patient’s prognosis and what treatment options are appropriate to recommend • ensuring familiarity with the patient’s history and current condition (this may include discussion with key community care providers), their family structure, and cultural needs and preferences • arranging adequate time for uninterrupted discussion • ensuring that patients have access to their regular communication aids • arranging for the appropriate people to be in attendance – ideally, this will include the patient; their substitute decision-maker, carers and family members, the most senior doctor available, the nurse responsible for the patient’s care and other members of the interdisciplinary team such as interpreters, aboriginal support workers, chaplains or social workers • ensuring that discussions can be held in an appropriately quiet and private environment. <p>1.6 The patient, substitute decision-maker, family and carers should be provided with written information about which clinician is responsible for leading and coordinating their care. Whenever possible, this clinician should be directly involved in discussions about the patient’s end-of-life care.</p> <p>1.7 Whenever possible, clinicians should work with patients, families and carers to ensure that key family members, substitute decision-makers and carers are present during end-of-life discussions. This will ensure that consistent messages are given about treatment options, their likelihood of success, risks and prognosis. This is particularly important when patients are in the process of transition to the terminal phase, as conflict and uncertainty among family and carers can result in avoidable suffering.</p>

Relevant standard	Relevant action (where available) against Essential Component 5
	<p>1.11 Clinicians should check that patients, families and carers have been provided with sufficient support to make decisions. This includes support for patients, substitute decisionmakers, families and carers who have communication difficulties associated with cultural and linguistic diversity, or decision-making difficulties associated with disability, mental illness or cognitive impairment.</p> <p>Paediatric</p> <p>1.2 Clinicians should assess the capacity of the child to be involved in discussions and decision-making about their care. This capacity may change across the illness trajectory and as the child develops.</p> <p>1.4 Clinicians should be respectful, sensitive and responsive to the preferences and needs of individual children and their families, regardless of aspects of identity such as culture, religious belief, gender or sexual preference.</p> <p>1.6 Whenever possible, clinicians should prepare for having conversations about end-of-life care. Necessary preparation may include:</p> <ul style="list-style-type: none"> • reaching consensus among all teams involved in the child’s care about the prognosis and what treatment options are appropriate to recommend • ensuring familiarity with the child’s history and current condition (this may include discussion with key community care providers), their family structure, and cultural needs and preferences arranging adequate time for uninterrupted discussion • arranging adequate time and an appropriate physical environment for uninterrupted discussion • ensuring that the child and their parents have access to their regular communication aids • arranging for the appropriate people to be in attendance – depending on the child and parents’ individual preferences, this may include the child, their parents and family, the most senior doctor available, the nurse responsible for the child’s care and other members of the interdisciplinary team such as interpreters, Aboriginal support workers, child life therapists, chaplains or social workers • ensuring that discussions can be held in an appropriately quiet and private environment. <p>1.7 Clinicians should provide parents with written information about which clinician is responsible for leading and coordinating their child’s care. Whenever possible, this clinician should be directly involved in discussions about the child’s end-of-life care.</p>

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	<p>1.8 Clinicians should assist parents in communicating information to their child about the outcomes of discussions about end-of-life care.</p> <p>1.9 Clinicians should work with parents to ensure that key family members are present during end-of-life discussions or that appropriate follow-up meetings are offered. This will ensure that consistent messages are given about treatment options, their likelihood of success, risks and prognosis.</p> <p>1.10 In some cultures, mainstream assumptions about death and dying may not be correct. Clinicians should work with parents to identify as early as possible any culturally appropriate decision-makers to include in discussions about end-of-life care.</p> <p>1.13 Clinicians should check that the child and their parents have been provided with sufficient support to make decisions. This includes support for children and parents who have communication difficulties, cultural, spiritual or linguistic needs, or decision-making difficulties associated with disability, mental illness or cognitive impairment.</p>
<p>3. Goals of care Clear goals improve the quality of end-of-life care</p>	<p>Adult</p> <p>3.1 The psychosocial, cultural and spiritual needs of patients, families and carers should be assessed, and care should be provided in accordance with their wishes, values and needs. Support should be offered for patients, carers and families who wish to include cultural or religious practices in their care, such as particular foods, singing, ceremonies or healing.</p> <p>3.4 Unnecessary burdens associated with medical treatment should be avoided for dying patients. For example, non-beneficial or unwanted observations, surgical interventions, investigations, medications and treatments should not be prescribed or administered. Where a clinician feels pressured – by the patient, family, carer or another health professional – to provide such interventions, they should seek advice and support from an experienced colleague who is able to assist with a second opinion and sensitive discussion.</p> <p>3.5 The goals of care, the treatment plan and any limitations of medical treatment for the current admission should be appropriately discussed with the patient, and their substitute decision-maker, family and carers, and clearly documented in the patient clinical record by the treating medical officer or team. This should occur as early as possible in the patient’s admission to the health service, and the information should be routinely reviewed and updated throughout the admission.</p>

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	<p>3.6 The rationale for medical decisions to discontinue or withhold nonbeneficial observations, investigations or treatments should be clearly communicated with the patient, family and carers.</p> <p>3.7 The goals of care, treatment plan and any limitations of medical treatment should be revisited with the patient, substitute decision-maker, family and carers when significant changes in the patient’s condition or circumstances occur – for example, when the patient is in transition to the terminal phase. The patient, substitute decision-maker, family and carers should be empowered to request further discussion and a review of the plan at any time.</p> <p>3.10 All patients, including people with chronic or severe mental illness, intellectual disability or cognitive impairment, have the right to adequate pain relief and symptom control, and to the prevention and relief of suffering. Physical symptoms should be managed in alignment with the patient’s wishes, and treatment reviewed regularly. If communication problems make it difficult to ascertain the patient’s wishes or needs, it is important to seek advice from, and consult with, families and carers.</p> <p>Paediatric</p> <p>3.1 Clinicians should assess the psychosocial, cultural and spiritual needs of the child. Support should be offered for families who wish to include cultural or religious practices in the care of the child, such as particular foods, singing, ceremonies or healing.</p> <p>3.3 Clinicians should avoid unnecessary burdens associated with medical treatment for dying children. For example, non-beneficial observations, surgical interventions, investigations, medications and treatments should not be prescribed or administered. Where a clinician feels pressured – by the family or another health professional – to provide such interventions, they should seek advice and support from an experienced colleague who is able to assist with a second opinion and sensitive discussion or seek legal or clinical ethics advice where appropriate.</p> <p>3.5 Clinicians should revisit the goals of care, treatment plan and any limitations of medical treatment when significant changes in the child’s condition, preferences, or circumstances occur – for example, when the child is in transition to the terminal phase.</p> <p>3.6 Clinicians should clearly communicate the rationale for medical decisions to discontinue or withhold non-beneficial observations, investigations or treatments to the parents and, where appropriate, the child.</p> <p>3.10 Clinicians should support dying children to receive oral food and fluids for as long as they wish.</p>

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	<p>3.11 Clinicians should manage physical symptoms in alignment with the child and their parents' wishes, and review treatment regularly.</p> <p>3.14 Clinicians should support children to play, learn and socialise with friends. For adolescents this could include a boyfriend or girlfriend.</p>
<p>4. Using triggers Triggers identify when patients/children need end-of-life care</p>	<p>Adult</p> <p>4.2 A critically important trigger for assessment, discussion and consideration of referral to specialist palliative care is when the patient, family members, carers or other members of the interdisciplinary team request palliative care, or express concern or worry that the patient is dying or has unmet end-of-life care needs.</p> <p>4.3 Clinicians should consider other useful triggers for recognition and review of patients who may benefit from end-of-life care interventions. Such triggers might be derived from condition-specific mortality risk prediction tools or from critical events, such as:</p> <ul style="list-style-type: none"> • diagnosis of life-limiting conditions • poor or incomplete response to medical treatment, continued deterioration despite medical treatment or development of new clinical problems during inpatient admission. <p>Paediatric</p> <p>4.2 Clinicians should consider other useful triggers for recognition and review of children who may benefit from end-of-life care interventions. Such triggers might be derived from condition-specific mortality risk prediction tools or from critical events, such as:</p> <ul style="list-style-type: none"> • presentation with life-threatening trauma or disease • diagnosis of life-limiting conditions • poor or incomplete response to medical treatment, continued deterioration despite medical treatment, or development of new clinical problems during inpatient admission • repeated calls to the rapid response team, particularly if the child has been admitted for more than one week • multi-system comorbidities (cardiovascular, pulmonary, endocrine, etc.) • maximal medical therapies already in place • decline in the child's condition, or a clinical determination that they will not benefit from interventions such as surgery, dialysis or treatment in intensive care • unexpected or prolonged recent admissions to hospital for exacerbation of a life-limiting chronic condition • the child, parents and family members, or other members of the interdisciplinary team requesting palliative care or expressing concern or worry that the child is dying or has unmet end-of-life care needs.

Relevant standard	Relevant action (where available) against Essential Component 5
<p>5. Responding to concerns Clinicians get help to rapidly respond to patient/child suffering</p>	<p>Adult</p> <p>5.1 The criteria and processes for accessing help for end-of-life issues that are causing concern should be clearly defined in relevant policies and procedures.</p> <p>5.2 The patient, substitute decision-maker, family and carers should understand the triggers and process for requesting an urgent review, and the process for responding to their request.</p> <p>Paediatric</p> <p>5.2 Clinicians should provide information to the parents and the child so that they understand how to request an urgent review if they are concerned about their child's care, and the process for responding to their request.</p> <p>5.3 Clinicians should conduct an interdisciplinary review of the goals of care and the treatment plan when concerns are raised that a child has unmet end-of-life care needs including physical, psychosocial or spiritual distress. Plan and document required follow-up and ongoing communication with the child, parents, and other clinicians.</p> <p>5.5 When sudden or unexpected deterioration in the child's condition occurs, clinicians should inform the parents and the interdisciplinary team with primary responsibility for the child's care as soon as possible.</p>
<p>10. Supporting systems Systems align with NSQHS Standards to improve outcomes</p>	<p>Adult</p> <p>10.1 Systems should be in place to support clinicians to work with patients, families and carers to receive, prepare, review or update advance care plans and directives, according to the wishes of the patient. These systems should align with NSQHS <i>Standard 1 (Governance for safety and quality in health service organisations)</i> and <i>Standard 9 (Recognising and responding to clinical deterioration in acute health care)</i>.</p> <p>10.2 Systems should be in place to provide clinicians with access to essential palliative medications 24 hours a day, seven days a week. These systems should align with NSQHS <i>Standard 4 (Medication safety)</i>.</p> <p>Paediatric</p> <p>10.1 The health service organisation should ensure that systems are in place to support clinicians to work with children and their families to receive, prepare, review and/or update age-appropriate advance care plans, according to the wishes of the child and their family. These systems should align with NSQHS <i>Standard 1 (Governance for safety and quality in health service organisations)</i> and <i>Standard 9 (Recognising and responding to clinical deterioration in acute health care)</i>.</p>

Relevant standard	Relevant action (where available) against Essential Component 5
	10.2 The health service organisation should ensure that systems are in place to provide clinicians with access to essential palliative medications 24 hours a day, seven days a week. These systems should align with NSQHS <i>Standard 4 (Medication safety)</i> .

Standards for general practice (RACGP), 5th Edition

Relevant standard	Relevant action (where available) against Essential Component 5
Core Standard 1: Communication and patient participation	1.3 Informed patient decisions.
GP Standard 2.1: Continuous and comprehensive care	GP2.1 b. Our practice provides continuity of care and comprehensive care.

Aged Care Quality Standards (Australia)

Relevant standard	Relevant action (where available) against Essential Component 5
1. Consumer dignity and choice	
2. Ongoing assessment and planning with consumers	
3. Personal care and clinical care	
6. Feedback and complaints	

End-of-life and Palliative Care Framework (NSW Health)

Relevant standard	Relevant action (where available) against Essential Component 5
1. Care is person-centred	Care should be based on the unique, holistic needs and preferences of the person receiving care. It should respect their preferences and their dignity. The individual, their families and carers are equal partners in the decisions relating to their care and treatment. Provision of care should be based on assessed need and be flexible in response to the person's changing needs and preferences.
2. There is recognition and support for families and carers	Families and carers play a pivotal role in the end-of-life and palliative care service system. It is essential their role is recognised, valued, and supported. Health services should support families and carers to be involved in planning and providing care, and to access the services they need to carry out this role.

Clinical Principles for End-of-life and Palliative Care (NSW Health)

Relevant standard	Relevant action (where available) against Essential Component 5
<p>Key action 3: Comprehensive assessment</p>	<p>Comprehensive assessment enables the evaluation of physical, environmental, social, cultural, emotional, psychosocial and spiritual needs. Comprehensive assessment can also determine the need for inclusion of an Aboriginal liaison officer or palliative care Aboriginal health worker, as required, or the use of the Health Care Interpreter Service.</p> <p>Action: Processes are in place to ensure comprehensive, culturally appropriate and holistic assessment is undertaken for all people identified as having end-of-life and/or palliative care needs.</p>
<p>Key action 4: Care planning</p>	<p>Care planning establishes the care goals and needs of the person, their family and carers. Individuals should be involved as much as possible in their care planning, to ensure their preferences and needs are reflected. Recognising variation in community and cultural preferences and expectations of care will help to support a person-centred approach for all. Multicultural health and Aboriginal health professionals can provide cultural guidance and support and, where applicable, be part of the multidisciplinary team.</p> <p>Action: Processes are in place to ensure that following comprehensive assessment of people with life-limiting illnesses, the person and their family/carers are actively involved in discussions and decisions regarding immediate care needs and advance care planning. Appropriate documentation must reflect these decisions.</p>
<p>Key action 7: 24/7 access to support</p>	<p>Access to support 24 hours, seven days a week (24/7) delivers timely appropriate care interventions and builds partnerships in decision-making with the person, their family and carers. Providing access to urgent clinical advice, medications (prescribing and supply) and support in the after-hours period will support people to be cared for in their preferred place and provide a point of contact for individuals, their family and carers.</p> <p>Action: Processes are in place to ensure all people receiving end-of-life and palliative care, as well as their family and carers, have access to appropriate support 24/7, and are informed of how to access this support. In the after-hours period, mechanisms are in place to provide urgent clinical advice and support for people receiving care at home. People have clear information and instructions about how to access this advice.</p>