Module 6

Last Hours of Living
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Abstract

Clinical competence, willingness to educate and calm, and empathic reassurance are critical to helping patients and families in the last hours of living. Management of feeding and hydration, changes in consciousness, delirium, pain, breathlessness, and secretions are common. Management principles are the same at home or in a health care institution. However, death in an institution requires accommodations to assure privacy, cultural observances, and communication that may not be customary. In anticipation of the event, inform the family and other professionals about what to do and what to expect. Care does not end until the family has been supported with their grief reactions and those with complicated grief have been helped to get care.

Key words

Bereavement, breathing changes, complicated grief, coroner, decreased appetite, decreased mobility, decreased perfusion, decreased thirst, delirium, expectations, eyes, fatigue, imminent death, moving the body, neurological dysfunction, normal grief, pain, rigor mortis, sphincter control, swallowing, tasks of caring

Objectives

After reviewing this module, oncologists and other members of the cancer care team will be able to:

- Assist families to prepare for the last hours of life.
- Assess and manage the pathophysiologic changes of dying.
- Pronounce a death and notify the family.

Clinical case on trigger tape

A.F. is a 79-year-old woman with metastatic breast cancer who is in her own home, cared for by her daughter with the help of the home hospice program. She develops aspiration pneumonia, and it is treated with oral antibiotics. Advance care planning indicates she does not want to go to the hospital under any circumstances, and oral antibiotics were an intermediate level of care. The patient and daughter agree that, if she gets better, she may have some quality of time left. But, if she doesn’t, A.F. says she is ready to go. Her physician makes a joint home visit with the home hospice nurse in order to assess changes in mental status and because it sounds like her daughter panicked and considered calling 911.
Introduction

Virtually all oncologists participate in the care of dying patients and their families during their careers. Of all patients who die, only a few (<10%) will die suddenly and unexpectedly. Most (>90%) will die after a long period of illness with gradual deterioration until an active dying phase at the end. Care provided during those last hours and days can have profound effects, not just on the patient, but on all who participate. At the very end of life, there is no second chance to get it right.

We will first summarize the physiological changes that occur as patients are dying and approaches to the management of associated symptoms. Then, we will discuss care at the time of death. Loss, grief, and bereavement are covered in EPEC-O Module 4: Loss, Grief, and Bereavement.

Most oncologists have little or no formal training to manage the dying process or death. Most have neither watched someone die nor provided direct care during the last hours of life.

Families usually have even less experience or knowledge. Based on media dramatization and vivid imaginations, most people have developed an exaggerated sense of what dying and death are like. However, with appropriate management, it is possible to provide smooth passage and comfort for the patient and all those who watch.

Preparing for the last hours of life

During the last hours of their lives, all patients require skilled care around the clock. This can be provided in any setting as long as the professional, family, and volunteer caregivers are appropriately prepared and supported throughout the process. The environment must allow family and friends access to their loved one around the clock without disturbing others and should be conducive to privacy and intimacy. Medications, equipment, and supplies need to be available in anticipation of problems, whether the patient is at home or in a health care institution. As the patient’s condition and the family’s ability to cope can change frequently, both must be reassessed regularly and the plan of care modified as needed. As changes can occur suddenly and unexpectedly, caregivers must be able to respond quickly. This is particularly important when the patient is at home, if unnecessary readmission is to be avoided.

If the last hours of a person’s life are to be as rewarding as possible, advance preparation and education of professional, family, and volunteer caregivers are essential, whether the patient is at home, in an acute care or skilled nursing facility, a hospice or palliative care unit, prison, etc. Everyone who participates must be aware of the patient’s health status, his or her goals for care and the parents’ goals, if the patient is a child, advance directives, and proxy for decision making. They should also be knowledgeable about the potential time course, signs and symptoms of the dying process, and their potential management. Help families to understand that what they see may be very different from the patient’s experience. If family members and caregivers feel confident, the experience
can provide a sense of final gift giving. For parents of a dying child, confidence can leave a sense of good parenting. If unprepared and unsupported, they may spend excessive energy worrying how to handle the next event. If things do not go as hoped for, family members may live with frustration, worry, fear, or guilt that they did something wrong or caused the patient’s death.

Establish in advance whether potential caregivers, including professionals who work in institutions, are skilled in caring for patients in the last hours of life. Do not assume that anyone, even a professional, knows how to perform basic tasks. Those who are inexperienced in this particular area will need specific training including, for instance, knowledge about body fluid precautions. Written materials can provide additional support to caregivers when experts are not present.

Although we often sense that death will either come quickly over minutes or be protracted over days to weeks, it is not possible to predict when death will occur with precision. Some patients may appear to wait for someone to visit, or for an important event such as a birthday or a special holiday, and then die soon afterward. Others experience unexplained improvements and live longer than expected. A few seem to ‘decide to die’ and do so very quickly, sometimes within minutes. While we may give families or professional caregivers a general idea of how long the patient might live, always advise them about the inherent unpredictability of the moment of death.

Physiologic changes and symptom management

There are a variety of physiological changes that occur in the last hours and days of life, and when the patient is actually dying. Each can be alarming if it is not understood. The most common issues are summarized here. To effectively manage each syndrome or symptom, oncologists need to have an understanding of its cause, underlying pathophysiology, and the appropriate pharmacology to use.

Fatigue/weakness

Weakness and fatigue usually increase as the patient approaches the time of death. It is likely that the patient will not be able to move around in the bed or raise his or her head.² As this is irreversible, weakness and fatigue need not be resisted and most treatment to alleviate it can be discontinued. Joints may become uncomfortable if they are not moved.³ Continuous pressure on the same area of skin, particularly over bony prominences, will increase the risk of skin ischemia and pain.⁴ As the patient approaches death, providing adequate cushioning on the bed will lessen the need for uncomfortable turning.
<table>
<thead>
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<td>Fatigue, weakness</td>
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<td>Weight loss, muscle &amp; fat, notable in temples</td>
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<td>Decreasing fluid intake, dehydration</td>
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<td>Peripheral edema due to hypoalbuminemia</td>
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<td>Peripheral and central cyanosis (bluing of extremities)</td>
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<td>Mottling of the skin (livedo reticularis)</td>
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<td>Venous pooling along dependent skin surfaces</td>
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<td>Dark urine</td>
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<td>Purposeless, repetitious movements</td>
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<td>Abnormal breathing patterns – apnea, Cheyne-Stokes respirations, agonal breaths</td>
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<td>Gurgling</td>
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<td>Loss of sphincter control</td>
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<td>Maceration of skin</td>
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<td>Perineal candidiasis</td>
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<td>Pain</td>
<td>Facial grimacing</td>
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<td>Tension in forehead; between eyebrows</td>
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<td>Loss of ability to close eyes.</td>
<td>Eyelids not closed</td>
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<td>Whites of eyes showing (with or without pupils visible)</td>
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<td>Rare, unexpected events:</td>
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<td>Bursts of energy just before death occurs, the ‘golden glow’</td>
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<td></td>
<td>Aspiration, asphyxiation</td>
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**Cutaneous ischemia**

At the end of life, fatigue need not be resisted and most treatment to alleviate it can be discontinued (see EPEC-O Module 3: Symptoms). Patients who are too fatigued to move and have joint position fatigue may require passive movement of their joints every 1 to 2 hours. To minimize the risk of pressure ulcer formation, turn the patient from side to side every 1 to 1.5 hours and protect areas of bony prominence with hydrocolloid dressings and special supports. Do not use ‘donuts’ as they paradoxically worsen areas of breakdown. A draw sheet can assist caregivers to turn the patient and minimize pain and shearing forces to the skin. If turning is painful, consider a pressure-reducing surface (e.g., air mattress or airbed). As the patient approaches death, the need for turning lessens as the risk of skin breakdown becomes less important.

Intermittent massage before and after turning, particularly to areas of contact, can both be comforting and reduce the risk of skin breakdown by improving circulation and shifting edema. Avoid massaging areas of erythema or actual skin breakdown.

**Decreasing appetite/food intake, wasting**

Most patients lose their appetite. Unfortunately, families and professional caregivers may interpret cessation of eating as ‘giving in’ or ‘starving to death.’ Yet, studies demonstrate that parenteral or enteral feeding of patients near death neither improves symptom control nor lengthens life. Anorexia may be helpful as the resulting ketosis can lead to a sense of well being and diminish discomfort. Help direct families and caregivers to find alternate ways to express their need to provide appropriate physical care and emotional support to the patient.

Physicians can help families understand that loss of appetite is normal at this stage. Remind them that the patient is not hungry, that food either is not appealing or may be nauseating, that the patient would likely eat if he or she could, and that clenching of teeth may be the only way for the patient to express desires.

Whatever the degree of acceptance of these facts, it is important for physicians to help families and caregivers realize that food pushed upon the unwilling patient may cause problems such as aspiration and increase tensions (see EPEC-O Module 11: Withdrawing Nutrition, Hydration). Above all, help them to find alternate ways to provide appropriate physical care and emotional support to the patient so that they can continue to participate and feel valued during the dying process.

**Decreasing fluid intake, dehydration**

Most patients stop drinking. This may heighten onlookers’ distress as they worry that the dehydrated patient will suffer, particularly if he or she becomes thirsty. Most experts feel that dehydration in the last hours of living does not cause distress and may stimulate endorphin release that promotes the patient’s sense of well-being. Low blood pressure or weak pulse is part of the dying process and not an indication of dehydration.
Patients who are not able to be upright do not get light-headed or dizzy. Patients with peripheral edema or ascites have excess body water and salt and are not dehydrated.

Decreased fluid intake usually heightens onlookers’ distress as they worry that the dehydrated patient will suffer, particularly if he or she becomes thirsty. As with feeding, families and professional caregivers will need support to understand that this is an expected event. It may help families to understand that most experts in the field feel that dehydration in the last hours of living does not cause distress and may stimulate endorphin release that adds to the patient’s sense of well-being. Low blood pressure or weak pulse is part of the dying process and not an indication of dehydration. Patients who are not able to move off the bed do not get lightheaded or dizzy. Patients with peripheral edema or ascites have excess body water and salt and are not dehydrated.

Parenteral fluids, either intravenously or subcutaneously using hypodermoclysis, are sometimes considered, particularly when the goal is to reverse delirium. However, parenteral fluids may have adverse effects that are not commonly considered. Intravenous lines can be cumbersome and difficult to maintain. Changing the site of the angiocatheter can be uncomfortable, particularly when the patient is cachectic or has no discernible veins. Excess parenteral fluids can lead to fluid overload with consequent peripheral or pulmonary edema, worsened breathlessness, cough, and orotracheobronchial secretions, particularly if there is significant hypoalbuminemia.

**Mucosal/conjunctival care**

To maintain patient comfort and minimize the sense of thirst, even in the face of dehydration, maintain moisture in mucosal membranes with meticulous oral, nasal, and conjunctival hygiene. Moisten and clean oral mucosa every 15 to 30 minutes with either baking soda mouthwash (1 teaspoon salt, 1 teaspoon baking soda, 1 quart tepid water) or an artificial saliva preparation to minimize the sense of thirst and avoid bad odors or tastes and painful cracking. Treat oral candidiasis with topical nystatin or systemic fluconazole if the patient is able to swallow. Coat lips and anterior nasal mucosa hourly with a thin layer of petroleum jelly to reduce evaporation. Avoid perfumed lip balms and swabs containing lemon and glycerin, as these can be both desiccating and irritating, particularly on open sores. If eyelids are not closed, moisten conjunctiva with an ophthalmic lubricating gel every 3 to 4 hours, or artificial tears or physiologic saline solution every 15 to 30 minutes to avoid painful dry eyes.

**Cardiac dysfunction, renal failure**

As cardiac output and intravascular volume decrease at the end of life, there will be evidence of diminished peripheral blood perfusion. Tachycardia, hypotension, peripheral cooling, peripheral and central cyanosis, and mottling of the skin (livedo reticularis) are normal. Venous blood may pool along dependent skin surfaces. Urine output falls as perfusion of the kidney diminishes. Oliguria or anuria is normal. Parenteral fluids will not reverse this circulatory shut down.
Neurologic dysfunction

The neurological changes associated with the dying process are the result of multiple concurrent irreversible factors. These changes may manifest themselves in 2 different patterns that have been described as the ‘two roads to death’ (see figure 1). Most patients follow the ‘usual road’ that presents as decreasing level of consciousness that leads to coma and death.

Decreasing level of consciousness

The majority of patients traverse the ‘usual road to death.’ They experience increasing drowsiness, sleep most if not all of the time, and eventually become unrousable. Absence of eyelash reflexes on physical examination indicates a profound level of coma equivalent to full anesthesia.

Communication with the unconscious patient

Families will frequently find that their decreasing ability to communicate is distressing. The last hours of life are the time when they most want to communicate with their loved one. As many clinicians have observed, the degree of family distress seems to be inversely related to the extent to which advance planning and preparation occurred. The time spent preparing families is likely to be very worthwhile.

While we do not know what unconscious patients can actually hear, extrapolation from data from the operating room and ‘near death’ experiences suggests that at times their awareness may be greater than their ability to respond. Given our inability to assess a dying patient’s comprehension and the distress that talking ‘over’ the patient may cause, it is prudent to presume that the unconscious patient hears everything. Advise families and professional caregivers to talk to the patient as if he or she were conscious.

Encourage families to create an environment that is familiar and pleasant. Surround the patient with the people, children, pets, things, music, and sounds that he or she would like. Include the patient in everyday conversations. Encourage family to say the things they need to say. At times, it may seem that a patient may be waiting for permission to die. If this is the case, encourage family members to give the patient permission to ‘let go’ and die in a manner that feels most comfortable to them. The physician or other caregivers might suggest to family members other words like:

- I know that you are dying, please do so when you are ready.
- I love you. I will miss you. I will never forget you. Please do what you need to do when you are ready.
- Mommy and Daddy love you. We will miss you, but we will be OK.

As touch can heighten communication, encourage family members to show affection in ways they are used to. Let them know that it is okay to lie beside the patient in privacy to maintain as much intimacy as they feel comfortable with.

Terminal delirium

An agitated delirium may be the first sign to herald the ‘difficult road to death.’ It frequently presents as confusion, restlessness, and/or agitation, with or without day-night reversal. To the family and professional caregivers who do not understand it, agitated terminal delirium can be very distressing. Although previous care may have been excellent, if the delirium goes misdiagnosed or unmanaged, family members will likely
remember a horrible death ‘in terrible pain’ and cognitively impaired ‘because of the drugs’ and may worry that their own death will be the same.

In anticipation of the possibility of terminal delirium, educate and support family and professional caregivers to understand its causes, the finality and irreversibility of the situation, and approaches to its management. It is particularly important that all onlookers understand that what the patient experiences may be very different from what they see.

If the patient is not assessed to be imminently dying, it may be appropriate to evaluate and try to reverse treatable contributing factors. However, if the patient is in the last hours of his or her life, the condition is by definition irreversible. Focus on the management of the symptoms associated with the terminal delirium in order to settle the patient and the family.20

When moaning, groaning, and grimacing accompany the agitation and restlessness, they are frequently misinterpreted as physical pain.21 However, it is a myth that uncontrollable pain suddenly develops during the last hours of life when it has not previously been a problem. While a trial of opioids may be beneficial in the unconscious patient who is difficult to assess, physicians must remember that opioids may accumulate and add to delirium when renal clearance is poor.22,23 If the trial of increased opioids does not relieve the agitation or makes the delirium worse by increasing agitation or precipitating myoclonic jerks or seizures (rare), then pursue alternate therapies directed at suppressing the symptoms associated with delirium.

Benzodiazepines are used widely to treat terminal delirium as they are anxiolytics, amnesticsts, skeletal muscle relaxants, and antiepileptics.24 Common starting doses are:

- Lorazepam, 1–2 mg as an elixir, or the tablet pre-dissolved in 0.5–1.0 ml of water and administered against the buccal mucosa q 1 h PRN will settle most patients with 2 to 10 mg/24 h. It can then be given in divided doses, q 3-4 h, to keep the patient settled. For a few extremely agitated patients, high doses of lorazepam, 20 to 50+ mg/24 h, may be required.

- Midazolam 1–5 mg/hr SC or IV continuous infusion, preceded by repeated loading boluses of 0.5 mg q 15 min to effect, may be a rapidly effective alternative.

Benzodiazepines may paradoxically excite some patients.25 These patients require neuroleptic medications to control their delirium.

- Haloperidol 0.5–2.0 mg IV, SC, PR q 1 h PRN, titrate to settle, then nightly to q 6 h to maintain) given intravenously, subcutaneously, or rectally may be effective.26

- Chlorpromazine 10–25 mg PO, PR, IM nightly to q 6 h and titrate to effect intravenously or rectally is a more sedating alternative.27

Barbiturates or propofol have been suggested as alternatives.28,29
Seizures may be managed with high doses of benzodiazepines. Other antiepileptics such as phenytoin pr or IV, fosphenytoin SC, or phenobarbital, 60 to 120 mg pr, IV, or IM q 10 to 20 min PRN, may become necessary until control is established.

**Respiratory dysfunction**

Changes in a dying patient’s breathing pattern may be indicative of significant neurological compromise. Breaths may become very shallow and frequent with a diminishing tidal volume. Periods of apnea and/or Cheyne-Stokes pattern respirations may develop. Accessory respiratory muscle use may become prominent. A few (or many) last reflex breaths may signal death.

Families and professional caregivers frequently find changes in breathing patterns to be one of the most distressing signs of impending death. Many fear that the comatose patient will experience a sense of suffocation. Knowledge that the unresponsive patient may not be experiencing breathlessness or ‘suffocating,’ and may not benefit from oxygen which may actually prolong the dying process, may be very comforting. Low doses of opioids or benzodiazepines are appropriate to manage any perception of breathlessness (see EPEC-O Module 3: Symptoms).

Some oncologists express concern that the use of opioids or benzodiazepines for symptom control near the end of life will hasten death. Consequently, they feel they must invoke the ethical principle of double effect to justify treatment. While it is true that patients are more likely to receive higher doses of both opioids and sedatives as they get closer to death, there is no evidence that initiation of treatment, or increases in dose of opioids or sedatives, is associated with precipitation of death. In fact, the evidence suggests the opposite.

**Loss of ability to swallow**

Weakness and decreased neurological function frequently combine to impair the patient’s ability to swallow. The gag reflex and reflexive clearing of the oropharynx decline and secretions from the tracheobronchial tree accumulate. These conditions may become more prominent as the patient loses consciousness. Buildup of saliva and oropharyngeal secretions may lead to gurgling, crackling, or rattling sounds with each breath. Some have called this the ‘death rattle’ (a term frequently disconcerting to families and caregivers).

Once the patient is unable to swallow, cease oral intake. Warn families and professional caregivers of the risk of aspiration. Scopolamine or glycopyrrolate will effectively reduce the production of saliva and other secretions. Common starting doses are:

- Scopolamine, 0.2–0.4 mg SC q 4 h, or
  Scopolamine, 1–3 transdermal patches q 72 h, or
  Scopolamine, 0.1–1.0 mg/hr by continuous IV or SC infusion
• Glycopyrrolate, 0.2 mg SC q 4 to 6 h
  Glycopyrrolate, 0.4–1.2 mg daily by continuous IV or SC

They will minimize or eliminate the gurgling and crackling sounds, and may be used prophylactically in the unconscious dying patient. Anecdote suggests that the earlier treatment is initiated, the better it works, as larger amounts of secretions in the upper aerodigestive tract are more difficult to eliminate. However, premature use in the patient who is still alert may lead to unacceptable drying of oral and pharyngeal mucosa. While atropine may be equally effective, it has an increased risk of producing undesired cardiac and/or CNS excitation.³⁷

If excessive fluid accumulates in the back of the throat and upper airways, it may need to be cleared by repositioning the patient or postural drainage. Turning the patient onto one side or a semiprone position may reduce gurgling. Lowering the head of the bed and raising the foot of the bed while the patient is in a semiprone position may cause fluids to move into the oropharynx, from which they can be easily removed. Do not maintain this position for more than a few minutes at a time, as stomach contents may also move unexpectedly.

Oropharyngeal suctioning is not recommended. It is frequently ineffective, as fluids are beyond the reach of the catheter, and may only stimulate an otherwise peaceful patient and distress family members who are watching.

**Loss of sphincter control**

Fatigue and loss of sphincter control in the last hours of life may lead to incontinence of urine and/or stool. Both can be very distressing to patients and family members, particularly if people are not warned in advance that these problems may arise. If they occur, attention needs to be paid to cleaning and skin care. A urinary catheter may minimize the need for frequent changing and cleaning, prevent skin breakdown, and reduce the demand on caregivers. However, it is not always necessary if urine flow is minimal and can be managed with absorbent pads or surfaces. If diarrhea is considerable and relentless, a rectal tube may be similarly effective.

**Pain**

While many people fear that pain will suddenly increase as the patient dies, there is no evidence to suggest this occurs.

Though difficult to assess, continuous pain in the semiconscious or obtunded patient may be associated with grimacing and continuous facial tension, particularly across the forehead and between the eyebrows. The possibility of pain must also be considered when physiologic signs occur, such as transitory tachycardia that may signal distress. Do not over-diagnose pain when fleeting forehead tension comes and goes with movement or mental activity, eg, dreams or hallucinations. Do not confuse pain with the restlessness, agitation, moaning, and groaning that accompany terminal delirium. If the diagnosis is
unclear, a trial of a higher dose of opioid may be necessary to judge whether pain is driving the observed behaviors.

Knowledge of opioid pharmacology becomes critical during the last hours of life. The liver conjugates codeine, morphine, oxycodone, and hydromorphone into glucuronides. Some of their metabolites remain active as analgesics until they are renally cleared, particularly morphine. As dying patients experience diminished hepatic function and renal perfusion, and usually become oliguric or anuric, routine dosing or continuous infusions of morphine may lead to increased serum concentrations of active metabolites, toxicity, and an increased risk of terminal delirium. To minimize this risk, discontinue routine dosing or continuous infusions of morphine when urine output and renal clearance stop. Titrate morphine breakthrough (rescue) doses to manage expressions suggestive of continuous pain. Consider the use of alternative opioids with inactive metabolites such as fentanyl or hydromorphone.

**Loss of ability to close eyes**

Eyes that remain open can be distressing to onlookers unless the condition is understood. Advanced wasting leads to loss of the retro-orbital fat pad, and the orbit falls posteriorly within the orbital socket. As eyelids are of insufficient length to both extend the additional distance backward and cover the conjunctiva, they may not be able to fully appose. This may leave some conjunctiva exposed even when the patient is sleeping. If conjunctiva remains exposed, maintain moisture by using ophthalmic lubricants, artificial tears, or physiological saline.

**Medications**

As patients approach death, reassess the need for each medication and minimize the number that the patient is taking. Leave only those medications to manage symptoms such as pain, breathlessness, excess secretions, and terminal delirium and reduce the risk of seizures. Choose the least invasive route of administration: the buccal mucosa or oral routes first, the subcutaneous or intravenous routes only if necessary, and the intramuscular route almost never.

**Dying in institutions**

The preceding discussion is relevant to patients dying in any setting (e.g., at home, in hospital, in nursing homes, other extended care facilities, jails, etc). However, a few remarks are warranted regarding the particular challenges of ensuring a comfortable death in an institution whose culture is not focused on end-of-life care.

When death is imminent, it is appropriate that patients remain with caregivers they know, rather than be transferred to another facility. Institutions can help by making the environment as home-like as possible. It is appropriate for the physician to order a private room where family can be present continuously and be undisturbed with the patient if
they so choose. The physician will want to cue the professional staff and encourage continuity of care plans across nursing shifts and changes in house staff.

Priorities and care plans at the end of life differ considerably from those for life-prolongation and cure. It is frequently challenging for physicians and other health care professionals to incorporate both into a busy hospital or skilled nursing facility. For this reason, specialized units where patients and families can be assured of the environment and the skilled care they need have been developed in many institutions.41,42

**When death occurs**

No matter how well families and professional caregivers are prepared, they may find the time of death to be challenging. Families, including children, and caregivers may have specific questions for the physician.

**Table 2: Signs that death has occurred**

<table>
<thead>
<tr>
<th>Description</th>
<th>Details</th>
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</thead>
<tbody>
<tr>
<td>The heart stops beating</td>
<td></td>
</tr>
<tr>
<td>Breathing stops</td>
<td></td>
</tr>
<tr>
<td>Pupils become fixed and dilated</td>
<td></td>
</tr>
<tr>
<td>Body color becomes pale and waxesen as blood settles</td>
<td></td>
</tr>
<tr>
<td>Body temperature drops</td>
<td></td>
</tr>
<tr>
<td>Muscles and sphincters relax (muscles stiffen 4-6 hours after death)</td>
<td>as rigor mortis sets in</td>
</tr>
<tr>
<td>Urine and stool may be released</td>
<td></td>
</tr>
<tr>
<td>Eyes may remain open</td>
<td></td>
</tr>
<tr>
<td>The jaw can fall open</td>
<td></td>
</tr>
<tr>
<td>Observers may hear the trickling of fluids internally, even after death</td>
<td></td>
</tr>
</tbody>
</table>

Basic information about death may be appropriate, e.g., the heart stops beating; breathing stops; pupils become fixed; body color becomes pale and waxy as blood settles; body temperature drops; muscles and sphincters relax, urine and stool may be released; eyes may remain open; the jaw can fall open; and observers may hear the trickling of fluids internally.43,44

There are no universally applicable ‘rules’ that govern what happens when the patient dies (in any setting). If the patient dies an expected death at home there is no need to call for emergency assistance. If a hospice program is involved, have the family call the hospice. If a hospice program is not involved, determine in advance who should be notified. Unless death is unexpected, or malice is suspected, involvement of the coroner’s office is usually not required (local regulations may vary).

When expected death occurs, the focus of care shifts from the patient to the family and those who provided care. Even though the loss has been anticipated for some time, no
one will know what it feels like until it actually occurs, and indeed it may take hours to
days to weeks or even months for each person to realize the full effect.

Many experts assert that the time spent with the body immediately after the death will
help people with acute grief. Those present, including caregivers, may need the
physician’s permission to spend the time to come to terms with the event and say their
good-byes. There is no need to rush, even in the hospital or other care facility. Encourage
those who need to touch, hold, and even kiss the person’s body as they feel most
comfortable (while maintaining universal body fluid precautions). As a visually peaceful
and accessible environment may facilitate the acute grieving process, a few moments
spent alone in the room positioning the patient’s body, disconnecting any lines and
machinery, removing catheters, and cleaning up any mess will allow the family closer
access to the patient’s body.

Spiritual advisors or other interdisciplinary team members may be instrumental in
orchestrating events to facilitate the experience of those present. Those who have not
been present for the death may benefit from listening to a recounting of how things went
leading up to the death and afterward. Grief reactions beyond cultural norms suggest a
risk of significant ongoing or delayed grief reactions.

When letting people know about the death, follow the guidelines for communicating bad
news. Try to avoid breaking unexpected news by telephone, as communicating in person
provides much greater opportunity for assessment and support. If additional visitors
arrive, spend a few moments to prepare them for what they are likely to see.

Once family members have had the time they need to deal with their acute grief reactions
and observe their customs and traditions, then preparations for burial or cremation and a
funeral or memorial service(s) can begin. Some family members may find it therapeutic
to help bathe and prepare the person’s body for transfer to the funeral home or the
hospital morgue. For many, such rituals will be their final act of direct caring.

Depending on local regulations and arrangements, some funeral directors will insist on
the completed death certificate being present before they pickup the body. All will
require a completed death certificate to proceed with any body preparation and
registration of the death. To avoid delaying the process, ensure that the physician who
will complete the certificate has ample warning that one will be required.

For many, moving the body is a major confrontation with the reality of the death. Some
family members will wish to witness the removal. Others will find it very difficult and
will prefer to be elsewhere. Once the body has been removed and family members are
settled, professional caregivers can offer to assist them with some of their immediate
tasks. They may notify other physicians and caregivers that the death has occurred so that
services can be stopped and equipment removed. Local regulations governing the
handling of medications and waste disposal after a death vary. When family members are
ready, professional caregivers can let the family know how to reach them, and then leave
them to have some privacy together.
Pronouncing death

In teaching hospitals, medical students and residents are typically called to ‘pronounce’ death. In non-teaching settings, the attending physician or nursing staff may be the ones to do it. When a patient dies at home with hospice care, it is usually a nurse who confirms the absence of vital signs. Although local regulations differ, if an expected death occurs at home without hospice care and the patient has a physician willing to sign a death certificate, then transportation to a hospital for a physician to confirm death may not be needed.

The phone call: “Please come, I think the patient has died.”

- Find out the circumstances of the death – expected or sudden? Is the family present? What is the patient’s age?

Preparation before you enter the room

- Confirm the details on the circumstances of death. Ask a nurse or other caregiver. Review the chart for important medical (length of illness, cause of death) and family issues (who is family?, faith?, is there a clergy contact?).
- Find out who has been called. Other physicians? The attending?
- Has an autopsy ever been requested? Do you see a value in requesting an autopsy?
- Has the subject of organ donation been broached? Has the Organ Donor Network been contacted?

In the room

- You may want to ask the nurse or chaplain to accompany you; he/she can give you support and introduce you to the family.
- Introduce yourself (including your relationship to the patient) to the family if they are present. Ask each person their name and relationship to the patient. Shake hands with each.
- Say something empathic: “I’m sorry for your loss...” or “This must be very difficult for you...”
- Explain what you are there to do. Tell the family they are welcome to stay if they wish, while you examine their loved one.
- Ask what questions the family has. If you cannot answer, contact someone who can.

The pronouncement

- Identify the patient. Use the hospital ID tag if available. Note the general appearance of the body.
- Test for response to verbal or tactile stimuli. Overtly painful stimuli are not required. Nipple or testicle twisting, or deep sternal pressure, are inappropriate.
• Listen for the absence of heart sounds; feel for the absence of carotid pulse.
• Look and listen for the absence of spontaneous respirations.
• Record the position of the pupils and the absence of pupillary light reflex.
• Record the time at which your assessment was completed.

Documentation in the medical record

• Called to pronounce (name); chart findings of physical examination.

• Note date and time of death; distinguish time family or others noted death from the time you confirmed the absence of vital signs. Note if family and attending physician were notified.

• Document if family declines or accepts autopsy; document if the coroner was notified.

Telephone notification

There will be cases where the people who need to know about the death are not present. In some cases, you may choose to tell someone by phone that the patient’s condition has ‘changed,’ and wait for them to come to the bedside in order to tell the news. Factors to consider in weighing this include: whether death was expected, the anticipated emotional reaction, if the person is alone, if the person is able to understand, distance, availability of transportation, and time of day. Inevitably, there are times when notification of death by telephone is unavoidable. If this is anticipated, prepare for it. Determine who should be called and in what fashion. Some families will prefer not to be awakened at night if there is an expected death.

1. Get the setting right. Determine the facts before you call. Find a quiet or private area with a telephone. Identify yourself and ask the identity of the person you are talking to and their relationship to the patient. Ask to speak to the person closest to the patient (ideally the health care proxy or the contact person indicated in the chart). Avoid responding to direct questions until you have verified the identity of the person to whom you are speaking. Ask if the contact person is alone. Do not give death notification to minor children.

2. Ask what the person understands about the patient’s condition with a phrase like, “What have the doctors told you about M’s condition?”

3. Provide a ‘warning shot’ like, “I’m afraid I have some bad news.”

4. Tell the news. Use clear, direct language without jargon like, “I’m sorry to have to give you this news, but M just died.” Avoid words like expired, passed away, and passed on. They are easily misinterpreted.

5. Respond to emotions with empathy. Most importantly, listen quietly to the person, including permitting enough time for the information to sink in. Elicit questions with a
phrase like, “What questions do you have?” Ascertained what support the person has. Ask if you can contact anyone for them. Consider other support through the person’s church, Red Cross, local police, or other service agencies if it is needed.

6. **Conclude with a plan.** If the family chooses to come to see the body, arrange to meet them personally. Provide contact information for the physician or official who can meet with them and/or make arrangements.

Immediately after the death, those who survive will need time to recover. A bereavement card or letter of condolence from the physician and attendance at the patient’s funeral may be appropriate. For many physicians and members of the professional team, encouraging follow-up visits from family members to assess the severity of their grief reactions to their recent loss, coping strategies, and provide support is a part of their professional duty of care. Professional members of the interdisciplinary team can also offer to assist family members to deal with outstanding practical matters, secure documents to redeem insurance, find legal counsel to execute the will, meet financial obligations, close the estate, etc. Bereavement care for the family is a standard part of hospice care in the US.

**Summary**

Clinical competence, willingness to educate and calm, and empathic reassurance are critical to helping patients and families in the last hours of living. For the majority of dying patients, predictable physiological changes occur. Management principles are the same at home or in a health care institution. However, death in an institution requires accommodations like assuring privacy, cultural observances and communication that may not be customary. In anticipation of the event, it helps to inform the family and other professionals about what to do and what to expect, including matters such as when rigor mortis sets in, how to call the funeral home, say goodbye, and move the body. Care does not end until the physician has helped the family with their grief reactions and helped those with complicated grief to get care. Care at the end of life is an important responsibility for every oncologist, and there is a body of knowledge to guide care.

**Key take-home points**

1. There is only one chance to get the management of the last hours right.
2. Patients usually need skilled care around the clock. The environment must allow family and friends ready access to their loved one in a setting conducive to privacy and intimacy.
3. Advance preparation and education of professionals, family, and volunteer caregivers are essential. They should also be knowledgeable about the potential time course, signs and symptoms of the dying process, and their potential management. Help
families to understand that what they see may be very different from the patient’s experience.

4. The physiologic changes of dying are complex. To control each symptom effectively, physicians need to have an understanding of its cause, underlying pathophysiology, and the appropriate pharmacology to use.

5. At the end of life, fatigue need not be resisted and most treatment to alleviate it can be discontinued.

6. Most patients lose their appetite and reduce food intake long before they reach the last hours of their lives. Anorexia may be protective, and the resulting ketosis can lead to a greater sense of well-being and diminish pain.

7. Most patients also reduce their fluid intake, or stop drinking entirely, long before they die. Dehydration in the last hours of living does not cause distress and may stimulate endorphin release that adds to the patient’s sense of well-being.

8. Maintain moisture in mucosal membranes with meticulous oral, lip, nasal, and conjunctival hygiene and lubrication.

9. The majority of patients experience increasing drowsiness, sleep most if not all of the time, and eventually become unarousable. Absence of eyelash reflexes indicates a profound level of coma equivalent to full anesthesia.

10. Presume that the unconscious patient hears everything.

11. Moaning, groaning, and grimacing accompanying agitation and restlessness are frequently misinterpreted as pain. Terminal delirium may be occurring. While a trial of opioids may be beneficial in the unconscious patient who is difficult to assess, benzodiazepines or sedating neuroleptics may be needed to manage terminal delirium. Benzodiazepines may cause paradoxical exciting effects; these patients require neuroleptic medications to control their delirium.

12. Diminished hepatic function and renal perfusion may change pharmacology of chronically administered medications.

13. Secretions from the tracheobronchial tree frequently accumulate. Scopolamine or glycopyrrolate will effectively reduce the production of saliva and other secretions.

14. Institutional dying presents particular challenges. Priorities and care plans at the very end of life differ from those for life prolongation and cure.

15. Planning discussions should cover personal, cultural, and religious traditions, rites, and rituals that may dictate how prayers are to be conducted, how a person’s body is to be handled after death, and when/how the body can be moved.

16. When expected death occurs, the focus of care shifts to the family and those who provided care.

17. Attend to acute grief reactions, especially when the body is moved.
Pearls

1. Use only essential medications. Stop routine dosing and continue to offer opioids ‘as needed.’ Accumulating serum concentrations of active drug and metabolites may lead to toxicity and terminal delirium.

2. Know the signs of the dying process.

Pitfalls

1. Maintaining parenteral fluids. They may have adverse effects that are not commonly considered.

2. Oropharyngeal suctioning. It is likely to be ineffective at clearing secretions, yet very effective at stimulating a gag, cough, or vomiting.

3. Removal of the body insensitively or too soon. This can be more distressing for families than the moment of death.

References


    Throughout its trajectory, Palliative Care can relieve the intense, broad suffering of persons living with the disease.


    Patients in a persistent vegetative state or coma, or suffering from dementia, do not experience pain and suffering when food and water are withdrawn.

In patients with advanced dementia, there is no data to suggest that tube feeding prevents aspiration pneumonia, prolongs survival, reduces the risk of pressure sores or infections, improves function, or provides palliation. Risks are substantial. For severely demented patients the practice should be discouraged on clinical grounds.


Of the 32 patients terminally ill with cancer monitored during the 12 months of study, 20 patients (63%) never experienced any hunger, while 11 patients (34%) had symptoms only initially. Similarly, 20 patients (62%) experienced either no thirst or thirst only initially during their terminal illness. In all patients, symptoms of hunger, thirst, and dry mouth could be alleviated, usually with small amounts of food, fluids, and/or by the application of ice chips and lubrication to the lips.


This editorial supports the contention that administration of fluids to a dehydrated dying patient should be based on the clinical manifestations of water and electrolyte disorders and of the potential benefits of treatment. Does dehydration cause symptoms? Are these symptoms distressing to the patient? Do fluid and electrolyte administration and other treatments help make dying patients comfortable?


82 subjects with cancer. All subjects died without artificial fluid therapy after median 2 days in study. Over 50% of subjects had a serum osmolality less than 295 mOsm/kg. No statistically significant relationship was demonstrated between the level of hydration and respiratory tract secretions. No statistically significant association was found between level of hydration and thirst or dry mouth.


Of 19 dying patients with cancer receiving IV hydration ranging from 500 mL to 3000 mL, six experienced mild thirst, eight moderate thirst, and four severe thirst.


Health professionals' attitudes to dehydration in dying cancer patients and the special ethical problems associated with the question are analyzed. Guidelines to facilitate the decision-making process related to the initiation of parenteral fluids in dying patients, and a discussion on the law as it pertains to the treatment of dehydration in such patients, are presented.


58 consecutive patients with advanced cancer received hypodermoclysis. A solution of two-thirds 5% dextrose and one-third saline was used in all cases at a rate of 20-100 cc/hr, with 750 units of hyaluronidase per liter, and 20-40 mEq KCl/L were administered in all patients. The mean duration of infusion and volume were 14 +/- 9 days and 1.3 +/- 0.8 L/day, respectively (range, 0.5-3 L/day). Toxicity consisted in local infection in two cases and bruising in two cases.

Co-Editor Dame Cicely Saunders, OM, DBE, FRCP (St Christopher's Hospice, London) is often regarded as the founder of the modern hospice movement. She provides an excellent overview chapter which includes the history of approaches to management of terminal cancer. Her colleague and co-editor, Nigel Sykes, MA, MRCGP (St Christopher's Hospice, London) is also a recognized leader in palliative care.


Delirium occurs commonly in the terminally ill and is associated with a poor prognosis. Management requires a clear understanding of what delirium is, how to assess it, investigations for reversible causes, and the medical treatments that are most useful.


Statistically significant improvements in mental state and nausea and vomiting occurred following a change from morphine to oxycodone.


The median PR dose of chlorpromazine was 25 mg every 4–12 hr. The median IV dose was 12.5 mg every 4–12 hr. Eighteen patients had complete relief and two had partial relief before death.

In this article, situations in which the nonlethal use of barbiturates may be indicated and situations in which their administration can be morally justified are discussed.


A low-dose infusion of the anesthetic agent propofol (Diprivan, Zenica), sedates patients who are not sedated by midazolam.


The incidence of pain, dyspnea, moist breathing, nausea and vomiting, confusion, restlessness, jerking and twitching, difficulty in swallowing, incontinence and retention of urine, sweating, moaning and groaning, and loss of consciousness was assessed in 200 patients. Many of the features can be attributed to organic brain disease consequent to metabolic disorder associated with multi-organ failure.


Current treatment recommendations for the most important symptoms arising in the endstage of neurological diseases is compiled. These recommendations include treatment of dyspnea, death rattle, restlessness, pain, thirst, depression, and others. A discussion of difficult decisions is included, e.g., the appropriate extent of fluid substitution or the ethical implications of sedation in the terminal phase.


Retrospective chart review of 237 consecutive patients who died in a specialist palliative care unit. Of these, 13% received sedatives for 7 days or more, while 56% commenced sedative use only in the last 48 hours of life. The groups receiving no sedation or sedation for less than 48 hours had the shortest survival from admission (mean, 14.3 and 14.2 days), whereas the 7-day sedation group survived for a mean of 36.6 days (P<.001).


Synopsis: An award-winning account by a noted surgeon describing in frank yet compassionate detail just what most of us are likely to face when the time comes. Dying has become such a taboo subject in modern society, most of us never have the honor of being with someone in their final days and moments.

Chapters cover different types of death, making clear the processes and choices that go along with each one. It addresses both medical and emotional realities of common conditions such as cancer, heart disease, AIDS, Alzheimer’s, severe trauma, and just plain wearing out.

Why read such a book? Taking away the fear of the unknown can bring courage and peace in the face of a difficult time. This book presents unpleasant facts in simple language that anyone can understand.

Be prepared to cry, since reading this book may make you experience feelings associated with people you love.

With a standardized approach and early intervention, retained secretions were controlled in only 50% of patients, highlighting the importance of explanation in easing relatives' distress.


100 consecutive admissions to an acute palliative care unit were assessed. To characterize family satisfaction, a survey was sent to the families of 240 patients who had been cared for on the inpatient unit.


The business plan and the process followed to develop the Palliative Care Program at the Cleveland Clinic Foundation (CCF) is presented. The benefits generated from this effort and the lessons learned that may be helpful to other program managers are described.


The Hospice Institute of the Florida Suncoast. Care at the Time of Death. Hospice Training Program. Largo, FL: The Hospice Institute of the Florida Suncoast; 1996.


Nurses are the professionals left to deal with dying patients' grief and anger.

This reviews the steps of the death pronouncement.


When telephone notification is unavoidable, there are suggested guidelines to follow. Doctors must weigh the benefits of truthfulness against the risk of potential harm resulting from abrupt disclosure of the bad news.


The primary care physician needs an explicit understanding of how to determine the cause and manner of death and should use succinct, clear language in completing the death certificate. When doubt exists or an external cause of death is a possibility, the coroner or the medical examiner is the appropriate public health official to contact.


Survivors require our care: informed by specific knowledge, educated experience, and a compassionate attitude.


This sounding board presents the important reasons for families and physicians, for the physicians’ attendance at a patient’s funeral.


This review adapted from the Education for Physicians on End-of-Life Care (EPEC) Curriculum describes the predictable course that occurs as well as approaches to management. Management principles are the same at home or in a health care institution.

Hallenbeck J. Palliative care in the final days of life: "they were expecting it at any time". JAMA. 2005 May 11;293(18):2265-71. [PMID: 15886382] [Full Text].