End-of-life care for patients with chronic disease: the need for a paradigm shift

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Death comes to all of us, and if we survive into our late 70s or 80s, progressive organ failure, often with multiple co-morbidities, often characterises the pathway towards the end of life. The care of patients with end-stage cardiac, renal or respiratory failure feature increasingly in the provision of health services, and the costs are immense particularly in the last year of life [1].

However, it is also increasingly apparent that our current model of care does not provide what is best for patients at end-of-life. No one clinical service is equipped to provide for the patient’s needs at all stages of their illness trajectory. Indeed, the philosophy of care and management priorities often differ between service providers. This leads to discontinuity of care (see Figure 1). The emergency department and medical teams are geared to dealing with acute deterioration: the model is predominantly curative or “patch up and mend”. Palliative care and hospice teams focus on “end-of-life” much more readily, but tend to operate in another domain, often separated from acute services not just philosophically but often geographically or by cost-centre! Because each of us operates in our separate silos, moving from a “curative” to an “end-of-life” management approach is difficult. Even where the diagnosis of dying has been embraced, our behaviours are more powerfully governed by the context in which we work. Often the default position is to continue as before, however inappropriate that may be.

A new model is required. Appropriate end-of-life care means less intensive, non-curative, symptom-relieving support in which preparing for death is seen as more important than clinging on to life. This means that “a good death” should be regarded as a quality outcome for all clinical services irrespective of where and by whom they are provided. It means striving to provide continuity of care at the end-of-life. It means that chronic disease management, palliative care, end-of-life care and terminal care are regarded as a continuum to which all health care providers contribute (see Figure 2). Whether in rest homes, primary care, emergency departments, medical wards or outpatient clinics, the “diagnosis of dying” should be entertained, sensitively communicated, and allowed to shape subsequent management.

In our own Unit, a very bad death made us realise that there was a significant gap between our intentions and what we actually delivered. Since that incident, we have been attempting to improve end-of-life care in the Respiratory Medicine service in Dunedin Hospital. But we recognise that the obstacles are considerable, not because of attitudes on the part of individuals, but because “the system” militates against it. We have adopted several practical
tools which can be applied to improving end-of-life care, but we realise that these have limited impact unless they are accepted across the wider organisation of a District Health Board. In isolation, progress is almost impossible. As well as specific tools, there is a need for strategic initiatives. The approach has to be “both ... and”.

The tools

An Advanced Care Plan provides the opportunity for patients, their family, and health care providers to enter into the territory of “end-of-life”. Importantly, it opens up conversations. In many cases it is liberating – from denial of the reality that a patient is experiencing and from fear of what might lie ahead. The New Zealand Advanced Care Planning (ACP) Co-operative has been established through the Minister of Health. Excellent guidelines on the principles and application of ACP have just been published [2,3]. Advanced Care Planning is not the prerogative of a single professional group - specialists, or GPs, or palliative care physicians.

In Dunedin, we have started a Respiratory Failure Supportive Care Clinic which includes, among other things, the opportunity to introduce the concept of ACP. The qualification for referral to the Clinic is the so-called “surprise question” i.e. would we be surprised if the patient were to die within the next year? Areas for discussion include the medical prognosis, the patient’s hopes and fears for the future, palliative treatments that are currently needed, as well as ACP i.e. treatments that would be acceptable and those that would be excessive or futile in the event of acute deterioration. A generic ACP needs to be modified for specific disease groups such as patients with respiratory failure, and we have recently done so.

Readers are welcome to use the Advanced Care Co-operative website to obtain a copy (http://acp.hiirc.org.nz).

Try as everyone might, there are still occasions when acute-on-chronic deterioration is too distressing to be managed at home and patients present to hospital. The context of deterioration needs to be urgently considered (is this an end-of-life or terminal event?). The concept of Ceiling of Care is relevant in this setting, and derives from the ACP. The aim is to provide guidance to admitting staff who do not know the patient, so that there is continuity with the patients’ previously expressed wishes, and/or limitations to their treatment are clear. We are currently working to have Ceiling of Care information electronically tagged to the patient’s NHI (see Figure 3), so that on admission, along with adverse drug reactions, the information is readily available. Of course patients may change their minds about how much intervention is desirable or appropriate – the approach cannot be rigid. But in our experience having the “ceiling of care” defined at the time of admission provides direction and security, particularly to nursing staff, as to how the patient is to be managed. In some centres, the Liverpool Care Pathway is also used, again providing a framework for appropriate in-patient management (and not abandonment) of the terminally ill.

There is also immense scope for improving end-of-life care in the patient’s home and in rest homes, and many in the primary care sector are working to this end. The introduction of ACP in rest homes
is an obvious need. But the tool cannot be applied in isolation. Developing the palliative care skills of community and practice nurses as well as rest home carers is an obvious area where resources need to be allocated. In Otago/Southland criteria for providing “Year of Care” support is going to be extended in 2012 to include end-of-life patients, identified using the “surprise question”. This is an example of how an appropriate philosophy of care and resource allocation can be integrated.

The strategies

Perhaps the most powerful incentive to improve end-of-life care is that this is what patients want [4, 5] and it’s something that we would want for ourselves. Attitudes to death and dying from cancer have been powerfully and positively influenced by the hospice movement. But the philosophy of care which has been nurtured in that particular setting now needs to be extended and integrated into institutions where “cure and mend” has historically been the over-riding objective. The time has come for “both … and” rather than “either … or”. I am grateful to the Southern DHB for a recent opportunity to advocate for these priorities as part of the “Putting the Patient First” Strategic Plan. The list of proposals currently being considered is in Table 1.

Conclusion

Patients at the end of life do not always want - and do not necessarily need – vigorous interventions but quality supportive care [6]. Quality improvement for such patients will be achieved not by straining indefinitely to extend life via acute medical services, nor by abandoning them when these fail. Adjusting what we do in the light of the diagnosis of dying, and managing the approach to death positively and meaningfully needs to be integrated into all clinical services, not just a few, so that a “good death” is included in what we mean by quality of life.

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About the author

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References

1. Chan W.C., Jackson G., Winnard D., Anderson P. Healthcare services funded by Counties Manukau District Health Board for people in the last year of life *NZ Med J* 2011; 124; 1-12


Table 1  Current goals for improving end-of-life care via DHB initiated strategic plans. The list is not exhaustive. These provide a unifying framework in which individual “tools” can be adopted and applied by different teams

| Goal #1: | to ensure that provision for end-of-life care and advanced care planning is included in the strategic and business plans for each clinical service operated by DHBs and PHOs |
| Goal #2: | to ensure that all clinical quality improvement initiatives within the DHB and PHO will address “quality of death” issues as much as they address “quality of life” |
| Goal #3: | to expedite nurse-led initiatives which will provide consistency in end-of-life care between the community and in hospital:  
- provide in-service training in palliative treatments for non-malignant diseases for community and practice nurses.  
- adoption and implementation of the Liverpool Care Pathway across all adult medical and surgical hospital wards |
| Goal #4: | to explore mechanisms whereby patients in community rest homes have an Advanced Care Plan (where appropriate), and that these plans are central to their management in acute situations |
Figure 1  The end-of-life trajectory for patients with end organ failure

Organ system failure: end-of-life trajectory

- High
- Low
- Occasions of discontinuity of care

Frequent admissions, self-care becomes difficult

2-5 years but death often seems “unexpected”

Death

Figure 2

The 20th century medical model

Acute care → Chronic disease management → Acute care → Chronic disease management → Acute care

The 21st century medical model

Chronic disease management

Acute care → Acute care → Acute care → End of life care

Terminal care
Figure 3

**CEILING OF CARE / RESUSCITATION PREFERENCES FOR ACUTE ON CHRONIC RESPIRATORY FAILURE**

**NAME** ....................................................... **NHI** .............................. **DATE** ..................................................

This patient has been attending the Respiratory Failure Support Clinic and/or has been an in-patient under the care of the Respiratory Service. The following Care Plan has been discussed and agreed with the patient, their family/whanau/carer, and has been confirmed and/or revised by the consultant specialist (electronically signed).

It should be used in the event of an admission to Dunedin Hospital with acute dyspnoea. Assuming that other diagnoses have been considered and excluded (e.g. pneumothorax), the patient’s acute respiratory distress should or should not include the following:

**SYMPTOM RELIEF:** e.g. LOW FLOW OXYGEN / OPIATES / INTRA-NASAL MIDAZOLAM / HALOPERIDOL

Select one or more as appropriate for the patient’s needs

- **ANTIBIOTICS**
  - YES / NO
- **PREDNISONE**
  - YES / NO
- **NON-INVASIVE VENTILATION (BIPAP)**
  - YES / NO
- **ICU / POSSIBLE MECHANICAL VENTILATION**
  - YES / NO
- **CPR IN THE EVENT OF CARDIO-RESPIRATORY ARREST**
  - YES / NO

Signed ....................................................... (Consultant) .................................. (Date)