The Right to Life Australia Inc.

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The Secretary
Select Committee on End of Life Choices in the ACT
Legislative Assembly for the ACT
GPO Box 1020
CANBERRA ACT 2601

Submission to Select Committee on End of Life Choices in the ACT

The ACT Assembly Select Committee on End of Life choices is seeking community views on issues involving end of life choices.

Although it may be claimed that these views should only emanate from ACT residents, nonetheless the ACT is in a unique position.

It is small and virtually could be claimed to be part of New South Wales situated as it is.

Legalising euthanasia in the ACT would mean that it would also affect residents in New South Wales.

It would be very hard to confine the practice to the ACT.

Clearly the idea to legalise patient killing in the ACT is a follow-on from the tragic passage of such legislation in Victoria in late 2017. The main reason that the idea of embracing the principle of ‘the life not worthy to be lived’ by any legislature is fraught with danger to the elderly and the vulnerable in the community.

Eventually the idea of bestowing on people a so-called right to die will lead to a duty to die.

We only have to look at those few places in the world which have legalised euthanasia to see how the numbers of killable patients have escalated along with the increase in reasons for allowing this.

Last official figures in the Netherlands showed that 431 patients had their lives ended without their request!
The practice of euthanasia in Belgium is even more out of control with a survey of nurses claiming that approximately 50% of euthanized patients were killed without their consent.

Most US states have voted against legalising euthanasia with only six having done so thus far.

Significantly in 2015, the British House of Commons voted overwhelmingly against legalising euthanasia in a vote of three to one (330:118)!

The Scottish and Welsh Parliaments also voted against similar legislation.

These UK legislatures were all influenced by what has been occurring in those few countries where it has been legalised. Professor Theo Boer of the Netherlands – a former advocate of legalised euthanasia addressed the House of Commons MPs warning them – don’t do it!

The two Minority Reports of the Victorian Legislative Council’s Committee of Inquiry into End of Life Choices were from Dr Daniel Mulino PhD MLC (ALP) and Mrs Inga Peulich MLC (Liberal) are included with my submission.

As Dr Mulino MLC summed up: “In only a very small minority of end of life situations are symptoms of pain unmanageable. He soberly concludes “In practice, euthanasia and assisted suicide are a disproportionate response that causes far more social harm than good”.

Margaret Tighe
PRESIDENT

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MINORITY REPORT

MINORITY REPORT

Mrs Inga Peulich MLC (LIB)
Member for South Eastern Metropolitan Region
As a member of the Legal and Social Issues Committee of the Legislative Council, I submit this minority report opposing several significant recommendations being made by the committee to support the establishment of a regime which allows physical assisted dying in Victoria, and in a narrow range of circumstances, to allow euthanasia.

Terms of Reference

The inquiry into End of Life Choices by the Victorian Legislative Council's Legal and Social Issues Committee was established on 7 May 2015. The Legislative Council agreed to a motion to inquire into, consider and report, no later than 31 May 2016, on the need for laws in Victoria to allow citizens to make informed decisions regarding their own end of life choices and, in particular, the Committee should —

(1) assess the practices currently being utilised within the medical community to assist a person to exercise their preferences for the way they want to manage their end of life, including the role of palliative care;

(2) review the current framework of legislation, proposed legislation and other relevant reports and materials in other Australian states and territories and overseas jurisdictions; and

(3) consider what type of legislative change may be required, including an examination of any federal laws that may impact such legislation.

Background

I opposed the establishment of this inquiry on the basis that Victorians had access to an outstanding program of palliative care available to terminally ill patients, and that the real intent of the motion was to work towards the establishment of a physician assisted dying regime. The Greens had unsuccessfully introduced this into the Victorian Upper House in 2008 which I opposed at the time on, personal, social and moral grounds.

From the outset, this inquiry was expected to deliver the recommendations which are contained in the report due to the strong predisposition of the committee to voluntary euthanasia, from the time of the inquiry's establishment as well as the loading of the inquiry with pro euthanasia submissions.

Overwhelming support for Palliative Care

I am strongly in support of recommendations which seek to improve the operations and funding of our well respected and effective palliative care system.

The management of pain, accessibility, and availability of palliative care, and its access by indigenous and multicultural Victorians are areas where ongoing improvement is occurring. This needs to be supported to ensure that pain is managed effectively and that no terminally ill person dies in pain.
Advance Care Planning

The notion of advance care planning to promote end of life choices being made by Victorians is in principle worthy of support.

I am however concerned that the content and the implementation of advanced care plans can be used as a vehicle for the promotion of physician assisted dying. I am also concerned that advanced care plans may be used to lock in advanced care directives which bind physicians and medical staff to executing wishes which may well be subject to change when a person is confronted by the prospect of death.

No person can predict the psychological disposition of a person, his or her will to live or die, or propensity to change their mind when confronted by death.

Furthermore, if one of the options in advance care planning is physician assisted dying or worse still, euthanasia (involving the administration of a lethal drug by a third person in instances where the patient cannot self-administer), then advance care plans become problematic. This is the reason why I oppose advance care plans if the legal framework in which advance care plans exist include physician assisted dying or a form of euthanasia.

Physician Assisted Dying and Euthanasia

The substantial part of this minority report arises from my strong opposition to physician assisted dying and any form of euthanasia.

Any proposal for a physician assisted dying regime or a form of euthanasia is not only a slippery slope, but people will die as a result of accident, error or misdiagnosis.

Any accidental loss of life – even the loss of one life, means such a regime cannot be justified, just as the loss of life, due to capital punishment, deliberate or due to a possible miscarriage of justice, cannot be justified and was the reason for its abolition.

Furthermore, is it possible to guarantee that a person facing death will not experience a last minute change of mind when staring death in the face?

Worse still, when unable to communicate their wishes on life and death issues, can we be confident that the actions of the medical profession or family members will be genuinely motivated by the best interests of the patient, their wishes and/ or their views? Can ending the life of a person who is unable to give informed consent ever be justified in a modern, democratic and multicultural society?

It is even more disturbing to consider the social impact of state sanctioned assisted suicide. What impact does the promotion of a physician assisted death regime have on our efforts to reduce suicide or youth suicide? This is a most serious social impact that has not been adequately considered by this or any other inquiry of which I am aware.

And lastly, how long will it be before those who have been campaigning for a broadening of eligibility for physician assisted dying to call for a broader set of criteria including those who suffer
experiential pain, mental health sufferers, persons who are inarticulate or even children and babies?

**Conclusion**

In conclusion, except those who are prepared to take a significant leap of faith to support calls for a physician assisted dying regime, a caring and compassionate society cannot support the taking of life when it is clearly impossible to have safeguards against errors of medical judgement, the accidental taking of human life, let alone the moral arguments which exist against state sanctioned death, no matter what language this is cloaked in.

Consensual physician assisted dying or even consensual euthanasia enshrined in advance care directives can never be a one way ticket to a destination without cases having been prone to manipulation and subjective interpretation. Nor should any scheme ever deny the patient concerned with every opportunity to change his or her mind right up to the very end, should such a regime ever be established.

Through the availability of palliative care, well-funded and accessible, delivered by caring, well trained and compassionate staff with all of the sensitivities required by patients suffering terminal illness and their families, terminal patients can be looked after compassionately.

End of life directives that exclude physician assisted death options, is the only safe and assured way of protecting and safeguarding against unintended deaths due to human error, accident, misdiagnosis or patients unable to express their views, or factor in a possible renewed desire to live at a critical point of any life taking regime.

Many of the arguments presented to the committee in favour of “dying with dignity” were based on a general commitment to principles of personal autonomy and rights.

However, a state legislated regime of physician assisted dying is not just an exercise of personal autonomy. The implications of such a regime for those who legislate, those who administer and those who may be victims of poor implementation, something not uncommon in society or our hospitals, are too severe and the recommendations of this report must be given closer scrutiny and rejected as a response of a compassionate society.

Pro euthanasia advocates claim that life is not devalued by permitting assisted suicide and euthanasia. Given that there can never be a guarantee that a regime will never result in a single accidental death is cold comfort who those who lose a loved one “accidentally” as a consequence of such a regime being in place.

The pressuring of vulnerable and sick people is another dimension which is a risk for which we cannot fully account – prospectively or retrospectively.

Also, is there a risk of creating a society where choosing death becomes an obligation for the patient so as to relieve family of the responsibility and cost of looking after an ill or disabled patient who is consuming resources associated with a continuation of life?
Will a pro death culture be created if we see the ill as being such a burden to society that their death serves to eliminate them or the elderly from our overcrowded hospitals or nursing homes?

Such a culture would undoubtedly be corrosive to the trust patients and families have in the medical profession and our health institutions.

The respect of personal autonomy enshrined in advanced care planning is a means of respecting the wishes of the individual patient who no longer wishes to continue receiving treatment for terminal illnesses.

Any regime which goes beyond that is no longer simply about personal autonomy and raises far too many issues for a compassionate society to contemplate.

Inga Peulich MLC
Member for the South Eastern Metropolitan Region
Minority Report by
Dr Daniel Mulino PhD MLC (ALP)
Member for Eastern Victoria Region
EXECUTIVE SUMMARY

The Majority Report does not establish the case for the legalisation of assisted suicide or euthanasia. It is mistaken in asserting that it is possible to fundamentally change medical practice in order to help the small minority of people who experience unendurable pain and that no one will be worse off as a result. The facts say otherwise.

The key arguments that will be developed in this report are that:

- It is possible to respect individual autonomy while not empowering health professionals to actively participate in acts of assisted suicide or euthanasia.
- Even if it can be argued that euthanasia or assisted suicide are justifiable in some instances, the negative consequences arising from legalisation far outweigh the benefits arising in that minority of cases.

This report acknowledges that extremely complex situations can arise at the end of life

Not all pain can be effectively managed, whether it be physical or psychological. In the face of unmanageable pain, some people express a clear desire to end their life. For most of us, witnessing great suffering by another person, particularly a loved one, prompts a strong desire to find a way to end that suffering. That is why many of the people who have a desire and willingness to give assistance to these wishes do so out of compassion and love.

The potential for very difficult situations to arise towards the end of life was reinforced by direct evidence provided to the Committee. This evidence was received in both written form and through testimony. All of the people who gave this evidence exhibited great bravery in telling their stories. This report acknowledges the importance of these peoples’ experiences.

It is also important to acknowledge that situations involving unendurable suffering are not limited to the end of life. They can arise as a result of non-terminal diseases, accidents or other changes in circumstances that have ongoing negative consequences on a person’s quality of life. Cases occurring near the end of life are generally focussed on in the euthanasia and assisted dying debate since these are widely seen to provide the strongest support for the case for intervention to give effect to a person’s wish to die.

It is possible to respect individual autonomy without supporting euthanasia or assisted suicide

For many, the importance of respecting individual autonomy underpins the case for euthanasia and assisted suicide. It is almost universally agreed that adults have a right to make informed choices about their medical treatment, including opting for a withdrawal of treatment. This report supports that proposition.
Euthanasia and assisted suicide are different. They are not simply a matter of whether to give effect to an individual’s choice about their own treatment. They necessarily involve third parties, usually medical practitioners, in acts that will intentionally result in death.

As such, acts of euthanasia and assisted suicide move from the private realm into the public realm. Public policy questions such as the risk of unintended consequences and proportionality are relevant and must be considered if regulatory intervention is to be justified.

**Euthanasia and assisted suicide are a last resort in only a tiny minority of cases**

While not all pain can be managed, it can be managed in the vast majority of cases and the proportion of cases in which pain can be managed is constantly increasing.

Experts in palliative care, oncology and related fields almost unanimously agreed that almost all symptoms arising from physical pain at the end-of-life can now be managed. The palliative care and oncology experts who gave evidence also stated that, over long careers, the number of people expressing a desire to have their life shortened was very small.

Even where there is an expressed desire to die, it is critically important to understand the nuances of such requests. Where the person making the request is experiencing depression or a mental illness, which is relatively common, there are usually other treatment options worth exploring. Holistic palliative care and other forms of assistance can often provide effective relief, even if not complete, and can often lead to a reversal in the expressed desire.

**In practice, euthanasia and assisted suicide are a disproportionate response that cause far more social harm than good**

The number of instances of euthanasia and assisted suicide is growing rapidly in all major jurisdictions where it is legal. This has been occurring for almost two decades in some jurisdictions, with no sign of abatement. The usage of euthanasia and assisted suicide in practice is far out of proportion to the situations that were originally used to justify the practice in these jurisdictions: namely, that small minority of cases where the symptoms of pain are unmanageable.

Moreover, the rapid growth in documented cases of euthanasia and assisted suicide probably materially understates the actual prevalence of the practice. There is a widespread failure of safeguards and procedures across jurisdictions, including low rates of reporting.

While legalisation was supposed to bring what was occurring in the shadows into the light, legalisation has simply pushed the boundary of what is legal out further and may have increased the amount of activity that occurs beyond the sight of regulators.

In countries with legalised euthanasia or assisted suicide, many vulnerable people are being placed in difficult situations in which they have to make irreversible, complex choices under a great deal of pressure. Evidence suggests that it is doubtful that safeguards are working as intended for such people.
CHAPTER 1 – FRAMING THE ISSUE

This report starts from the premise that adults have a right to control what happens to them. This includes a right to decide which medical treatments are administered to them and a right to have medical treatment fully withdrawn should a person decide that the burden of any treatment would outweigh the likely potential benefits.

A patient should be able to convey a desire to have treatment withdrawn either explicitly while conscious or through a clear statement of their preferences that could take effect should a person fall into an indefinite unconscious state. The importance of this right was reflected in the Medical Treatment Act 1988 and this report concurs with the right to refuse treatment or have treatment withdrawn as critical.

It is important to stress from the outset that, in cases of both euthanasia and assisted suicide, the ethical dimensions are inherently more complex since third parties are necessarily involved. It is not enough to assert a belief in the individual's right to autonomy. Any system of regulated euthanasia or assisted suicide must involve the active participation of members of the health profession and possibly also other regulatory arms of the state.

In addition to considering how to give effect to the wishes of the patient, the role of medical professionals and the health system more generally in actively bringing about the death of patients must be scrutinised in any discussion of active euthanasia or assisted suicide.

The role of professionals in a regulated euthanasia or assisted suicide regime could take a number of forms, including some or all of the following:

- One or more medical professionals providing approval before an individual is granted access to either euthanasia or assisted suicide.
- One or more medical or related professionals providing counselling before either euthanasia or assisted suicide is administered.
- One or more medical professionals (or a chemist, with approval) prescribing the drug/poison to be used in an assisted suicide.
- One or more medical professionals administering euthanasia.
- Medical professionals or others as deemed appropriate witnessing an assisted suicide.
- Medical professionals and others as deemed appropriate monitoring instances of euthanasia or assisted suicide and possibly hearing appeals at different steps of the regulated process.

Because regulating euthanasia and assisted suicide involves the state regulating how medical professionals can take active steps to bring about death, it is not just a question of individual autonomy.

Euthanasia and assisted suicide involve broader policy questions in relation to how the medical system can best work in the interests of individuals who are either terminally ill or in grave physical
or mental distress. Issues such as unintended consequences and whether procedural breaches can be managed lie at the heart of the issue.

In giving evidence to the House of Lords Select Committee inquiry into the Assisted Dying for the Terminally Ill Bill, Professor Alan Johnson, Emeritus Professor of Surgery at the University of Sheffield observed:

... the impression has been given that obeying patients' wishes is the overriding ethical imperative for doctors. Of course it is important, but it is not paramount. If it were, I would have done many unnecessary operations and some harmful operations in my time as a surgeon (Q 165).¹

He cited, as an example of the need to subordinate individual patient autonomy to the interests of the wider patient community on rare occasions, the refusal of doctors to prescribe antibiotics at a patient's request for relatively trivial conditions because to do so would "produce resistance which might have quite a serious effect on people further down the line".²

The Select Committee concluded that: "We are agreed that patient autonomy cannot be absolute and that there must be some limits set, in the interests of the wider community."³

In the context of euthanasia and assisted suicide, there is arguably a gain from giving effect to some patients' wishes to hasten their death. But this must be weighed against the potential for societal harm through a range of potential negative consequences, including: vulnerable people being pressured into euthanasia or assisted suicide; people having their death hastened without having given proper consent; and a gradual broadening of practices without transparent public consideration. As will be shown below, evidence from jurisdictions that have legalised euthanasia and assisted suicide shows that all of these are real risks.

The House of Lords Select Committee supported this framing of the issue:

... we cannot address the issue of personal autonomy in isolation and ... we must proceed to look at some of the 'real world' issues which have been raised and to try to assess the balance between greater personal choice for some people and increased potential harm for others ...⁴

What is the real choice that we face? It is important to acknowledge that neither current regulatory arrangements nor a world with legalised euthanasia/assisted suicide will be free from unendurable suffering. We are ultimately choosing between two systems in which some suffering will be difficult to treat or manage and in which regulation is difficult to perfectly enforce.

The Majority Report gave considerable weight to the evidence from the Coroners Court of Victoria. This evidence included details of a number of confronting situations in which people committed suicide, often in very distressing ways, in order to end suffering. What the Majority Report doesn’t

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¹ House of Lords, Assisted Dying for the Terminally Ill Bill – First Report, Select Committee on Assisted Dying for the Terminally Ill Bill, Session 2004-05, 3 March 2005, paragraph 45.
² Ibid, paragraph 45.
³ Ibid, paragraph 62.
⁴ Ibid, paragraph 63.
make clear is that many of the instances raised by the Coroners Court wouldn’t fall within the scope of its proposed regime. This highlights the difficult choice that we face in practice.

As Table 1 outlines, we do not face a choice between respecting autonomy and bringing dubious actions out of the shadows on the one hand versus an intransigent, knee-jerk protection of the status quo on the other. Rather, we face a choice between two imperfect situations. In neither will the wishes of all people to die be fully complied with. In neither will there be full and transparent compliance with the law.

Rather, the choice as to whether to reform or not should be guided by an informed judgement of the likely consequences for our overall treatment of people in vulnerable situations.

Table 1: Characteristics of current arrangements vs legalised euthanasia/assisted suicide

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<tr>
<th>Current arrangements</th>
<th>Legalised euthanasia or assisted suicide</th>
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<tr>
<td>Harrowing deaths exist - although in the vast majority of cases, physical pain can be managed effectively with modern pain relief and palliative care. These instances are a small and shrinking minority of overall deaths.</td>
<td>Harrowing deaths will still exist – although there will probably be less in the category of deaths explicitly allowed under the euthanasia/assisted dying regime. The degree to which there are less deaths involving unmanageable physical or psychological pain will depend upon the permissiveness of the scheme.</td>
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<td>Psychological pain that is difficult to treat exists – although treatment in this sphere is improving.</td>
<td>Psychological pain that is difficult to treat will continue to exist – unless a very permissive regime is introduced, such as in Belgium. Under such a permissive regime, the effectiveness of safeguards will be extremely difficult to ensure.</td>
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<td>The number of instances of euthanasia or assisted suicide will probably rise rapidly and this increase is likely to persist. In foreign jurisdictions with legalised euthanasia or assisted suicide, there has been a rapid and sustained growth in the number of deaths and in no jurisdiction is there evidence that this increase will slow down.</td>
<td>It is likely that the enforcement of safeguards will be difficult. In jurisdictions with legalised euthanasia or assisted suicide, there is systemic non-compliance with the law. This includes: • non-reporting of many instances of euthanasia/assisted suicide; • incomplete awareness by regulators of breaches of the law; and • incomplete enforcement by health regulators and law enforcement of breaches.</td>
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<td>The law is currently not fully enforced. This can take a number of forms, including: prosecutorial discretion and sentencing leniency, particularly in relation to cases where people are killed or assisted in their suicide by loved ones.</td>
<td>It is unclear if enforcement of the law will improve at all. In jurisdictions with euthanasia and assisted suicide, it is acknowledged by regulators that considerable activity occurs at the edges of the law and that they are powerless to monitor, let alone prevent, much of this activity.</td>
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CHAPTER 2 – ALMOST ALL CASES OF PAIN ARE CURRENTLY MANAGEABLE

In almost all cases where a patient experiences a great deal of pain, including terminal cases, the symptoms of that pain can be managed by current best practice in pain relief and palliative care. It is important to note that the capacity of health practitioners to manage the symptoms of pain is increasing over time.

2.1 Pain relief is effective in almost all cases

A number of expert practitioners in palliative care gave evidence to the Committee in relation to the frequency of cases in which it was possible to manage pain at the end of a patient’s life. While it is not possible to put a precise percentage on this proportion, the experts were unanimous in their assessment that it was rare that pain couldn’t be managed.

This included the following evidence:

**Associate Professor Daryl Jones, Austin Health:** “... in 20 years as a clinical practitioner, the number of patients I have seen who die to whom that applies is the overwhelming minority.”

**Associate Professor Peter Hunter, Alfred Health:** “We do know that in almost all patients, if we do a proper assessment and understand that the drugs are available, we can alleviate pain and we can do a good job of that.”

“I have never not been able to control anyone’s symptoms around pain once you have got them on the right treatment path, be it narcotic analgesia or infusion pumps et cetera, so I think that if anyone dies in pain, that is an absolute travesty because there is capacity to really manage that effectively in this day and age.”

Moreover, the proportion of patients for whom pain is manageable is rising. When asked if palliative care techniques had improved, Dr Michelle Gold, Director of the Palliative Care Unit at the Alfred Hospital stated that:

**Dr Michelle Gold:** There are always new medications available in that sense. We are having increasingly fruitful interactions with some of our interventional colleagues to provide pain relief with various procedures and interventions that are sometimes much better directed than the medications are and hopefully have fewer side effects. It is terrific that there are new techniques – new ways of delivering some of the old medication.

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1 Associate Professor Daryl Jones, Transcript of evidence, 5 August 2015, p24.
2 Associate Professor Peter Hunter, Geriatrician and Director of Aged Care, Alfred Health, Transcript of evidence, 15 October 2015, p55.
3 Dr Michelle Gold, Director of the Palliative Care Unit of the Alfred Hospital, Transcript of evidence, 23 July 2015, pp56-57.
2.2 Knowledge about pain relief options can provide considerable comfort

Many health practitioners who provided evidence to the Committee spoke of the considerable potential to provide patients with peace of mind by clearly communicating the pain relief options available to them. For example Ms Deidre Bidmade, Vice-President of the Warrnambool and District Community Hospice, stated that:

Ms Deidre Bidmade: ... in the last 10 years the biggest factor when somebody is dying is how I am going to die. Is my pain going to be out of control? These conversations before somebody is in their last phase of dying is so important, because if you allay those fears and reassess the fact that we have such wonderful drugs out there to ensure that people are not in pain and that we do have things like Niki syringe drivers, so that you can go home with that in situ and that can be titrated to the needs every day, if you can give people peace of mind that they are supported in the journey and that there is no need for acceleration of symptom, that in my experience has been most of the underlying issue.\(^8\)

This view was supported by Associate Professor William Silvester, President of the International Society of Advance Care Planning and End of Life Care:

Associate Professor William Silvester: I am amazed by the number of times that I have gone to walk away and some has called me back and said, ‘Look doctor, I didn’t tell you, but in fact I’ve been thinking about euthanasia, and now that you’ve been able to lift such a weight off my shoulders and I can now be sure and confident that I am going to get the care that I want at the end in the way that I want it, I can now concentrate on living as well as possible. I don’t have to think any longer about getting Nembutal or saving up all of my prescription tablets or whatever so that I can do something before I lose control, because now I can see that even when I reach a point where I no longer have control, I will still get what I want and I will not get what I do not want.\(^9\)

2.3 Very few people ask to be killed – especially if they are aware of care options

Dr Ranjana Srivastava recently wrote an internationally acclaimed book, “Tell me the truth”, dealing with the meaning of a good life and good death and the ethics of end-of-life interactions. She is an oncologist with considerable experience and adjunct associate professor in the Monash University Faculty of Medicine, Nursing and Health Sciences. Dr Srivastava gave evidence to the Committee in relation to the issue of how often people seek active intervention by a doctor to end their life:

Dr Srivastava: ... from 15 years of experience ... the most informed I can tell you is that in all my career there have been no more than two or three people at most, in the thousands of patients I have seen, who have said, ‘I have had enough. I want to die.’\(^10\)

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\(^8\) Ms Deidre Bidmade, Vice-President of the Warrnambool and District Community Hospice, *Transcript of evidence*, 30 July 2015, p27.

\(^9\) Associate Professor William Silvester, President of the International Society of Advance Care Planning and End of Life Care, *Transcript of evidence*, 23 July 2015, pp63-64.

\(^10\) Dr Ranjana Srivastava, Adjunct Associate Professor, Faculty of Medicine, Nursing and Health Sciences, Monash University and Oncologist, *Transcript of evidence*, 19 August 2015, p20.
This was corroborated by other evidence, including from Professor Peter Hudson, Director of Palliative Care at St Vincent’s Hospital:

**Professor Peter Hudson:** We have been collecting data in two large teaching hospitals for about 15 years now, and it shows that less than 1 per cent of patients referred to the palliative care consultation service actually request euthanasia.\(^{11}\)

Even where people ask to die or ask to be killed, the subtle nuances in what they are communicating can only be interpreted by an expert:

**Dr Natasha Michael:** One of the questions I am always asked is: do patients ask you to kill them? The answer is: yes. I do not use the term ‘request for euthanasia’; I use the term, ‘They express a desire for death’. Many might have heard me use this terminology or expression before.

People say one of three things to me: ‘I want to die’, ‘Let me die’ or ‘Kill me’. The people who are saying ‘I want to die’ are people who are saying: my suffering is so unbearable at this point in time, with my pain, my distress or the burden I am placing on my family, death has to be a better option. The answer to that is not injecting them with a drug or providing barbiturates; the answer to that is saying, ‘How can I help you with these factors that are making you feel that death has to be the better option?’ The ‘Let me dies’ are saying, ‘You know what, I have fought this fight. I have run the course. I am 85 — or I am 63 — I have had six lines of chemotherapy. I have had enough. Let me die’. The ‘Kill me’ is from a really, really small minority [emphasis added]. Most health professionals are not trained to unpick these things. They lump them together in a singular cohort, so you think of the danger you pose when people come to you and express a desire for death when actually what they are saying is, ‘My suffering is so unbearable that I just want you to help me’. The risk is phenomenal.\(^{12}\)

If you have an unskilled practitioner, you do not have the ability to distinguish these complex psychosocial phenomenons. The risk is too high ... I have seen too many people express a desire for death when actually all they are asking for is help.\(^{13}\)

To set up a system in which many GPs face a situation in which they have to interpret complex end-of-life situations will create significant risks particularly given that most GPs do not have training in this specific area of care and will face such situations rarely. As will be outlined below in detail, these risks are further compounded by the inadequacy of safeguards in practice.

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\(^{11}\) Professor Peter Hudson, Director, Centre for Palliative Care, St Vincent’s Hospital, *Transcript of evidence*, 24 February 2016, p4.

\(^{12}\) Dr Natasha Michael, Director, Palliative Care, Cabrini Health, *Transcript of evidence*, 16 September 2015, p9.

\(^{13}\) Dr Natasha Michael, Director, Palliative Care, Cabrini Health, *Transcript of evidence*, 16 September 2015, p10.
2.4 Conclusion: A very small number of cases are problematic under current arrangements

For the vast majority of people, modern pain relief and palliative care can manage physical pain right up to the end of life. If euthanasia or assisted suicide was legalised so as to deal with this type of situation, one would expect to see a small and relatively stable number of cases over time. As will be outlined in the following chapter, the evidence differs markedly from this in every major jurisdiction that has legalised euthanasia or assisted suicide.

Psychological pain is sometimes more difficult to manage. However, once psychological pain is used as a rationale for euthanasia or assisted suicide, difficulties arise as this type of pain is not particularly associated with end-of-life issues. It can therefore be used to justify the extension of euthanasia and assisted suicide regimes to non-terminal situations. Moreover, as will be argued in Chapter 4, allowing euthanasia or assisted suicide in cases involving profound psychological pain is problematic as the patient’s condition will likely impact on their capacity to provide consent.
CHAPTER 3 – THE FREQUENCY OF EUTHANASIA AND ASSISTED DYING IS RISING RAPIDLY

3.1 In all jurisdictions where euthanasia or assisted suicide is legal, there has been a sharp, sustained upward trend in the number of deaths – often over long periods of time

In all major jurisdictions where euthanasia or assisted suicide is legal, the number of cases has increased sharply ever since legalisation. This increase has been rapid and sustained. In no major jurisdiction is a plateau in numbers evident.

The majority of this increase is not due to the aging of society or to a growing awareness of end-of-life options. As will be outlined in Chapter 4, it is most likely to due to factors such as:

- An expansion in the categories of people eligible to opt for euthanasia or assisted dying either through formal legislative change or, sometimes less transparently, gradual changes in the interpretation and application of existing provisions.
- A “normalisation” of euthanasia or assisted dying in the medical system and across the broader culture.
- Systemic failures in safeguards.

The Majority Report sidesteps this issue. It provides no attempt to explain either why such persistent growth in cases is occurring or whether the risks associated with this trend can be managed.

3.2 A summary of empirical trends in the number of cases of euthanasia and assisted suicide across major jurisdictions

Over the next two pages, the number of cases of euthanasia and assisted suicide in the major jurisdictions that have legalised either procedure are set out.

The data from which the graphs and associated growth rates were derived was all obtained from public sources.

The period of data availability varies by jurisdiction, but in three instances, data is available for more than 10 years and in two jurisdictions for more than 15 years.
EUROPEAN JURISDICTIONS

Figures 1 to 3 contain the trend in the total number of cases of euthanasia and assisted dying in Belgium, the Netherlands and Switzerland over the period for which reliable data is available. Data is available for 12 years in the case of Belgium, 7 years in the case of Netherlands and 16 years in the case of Switzerland.

Figure 1: Belgium

Total Growth 2003-2015: 235 → 2,012
Compound annual growth rate: 19.6%

Figure 2: Netherlands

Total Growth 2008-2015: 2,331 → 5,516
Compound annual growth rate: 13.1%

Figure 3: Switzerland

Total Growth 1998-2014: 50 → 836
Compound annual growth rate: 19.2%
NORTH AMERICAN JURISDICTIONS

Figures 4 and 5 contain the trend in the total number of cases of euthanasia and assisted dying in Oregon and Washington State over the period for which reliable data is available. The data is available for 17 years in the case of Oregon and 5 years in the case of Washington State.

**Figure 4: Oregon**

*Graph showing the trend in total number of cases from 1998 to 2015 with a significant increase.*

- Total Growth 1998-2015: 16 → 132
- Compound annual growth rate: 13.2%

**Figure 5: Washington State**

*Graph showing the trend in total number of cases from 2009 to 2014 with a steady increase.*

- Total Growth 2009-2014: 64 → 170
- Compound annual growth rate: 21.6%

For Belgium, the data was compiled from the official statistics of the Federal Control and Evaluation Commission. These are reported to the Belgian legislature.¹⁴

For the Netherlands, the data is sourced from the Dutch Regional Euthanasia Review Committees.¹⁵

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¹⁵ The Netherlands has five regional euthanasia review committees. The review committees assess whether a doctor who has performed or assisted suicide, euthanasia, has complied with the due care criteria set out in the **Termination of Life on Request and Assisted Suicide (Review Procedures) Act**. An annual report is issued each year by the Dutch Regional Euthanasia Review Committees.
The statistics for Switzerland are derived from a number of sources. Swiss authorities published the first official statistics on assisted suicide in March 2012. According to the Federal Statistical Office, the number of Swiss residents who died from assisted suicide has increased continuously between 1998 and 2009. The Federal Statistical Office reported that, in 2009, approximately 300 assisted suicide deaths occurred (4.8 per 1000 deaths or 0.48% of all deaths) compared with fewer than 50 deaths in 1998. In 2014, the total number of assisted suicide deaths in Switzerland, including deaths at Dignitas and the Eternal Spirit clinic, was around 836.

Given the lack of data, Figure 3 is derived by linear interpolation between the data points provided by the Federal Statistical Office for 1998 (using 50 to be conservative in relation to growth rates) and around 300 in 2009. There is a further linear interpolation between the 2009 figure and 836 in 2014.

Recent media articles detailing the number of deaths at individual organisations corroborate this overall trend. It has been reported that the Swiss assisted suicide organisation (EXIT) helped 782 people end their lives in 2015, 199 more than the previous year (583 in 2014). For 2011, it has been independently reported that the right-to-die organisation EXIT assisted 416 deaths, up from approximately 348 deaths in 2010.

In Switzerland, a material number of deaths are from non-residents. The 2011 figure for non-resident deaths reported by the organisation Dignitas was 149. The Atlantic reported that, in the decade leading up to 2010, over 1,000 people had received assisted suicide at Dignitas.

The data for Oregon and Washington State is sourced from the relevant State Department of Health reports.

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19 EXIT reports jump in assisted suicide numbers, Swissinfo.ch, website, 2 March 2016. (Accessed 30 May 2016)


3.3 The sustained and long-term nature of growth

In all jurisdictions, the increase in the total number of cases has been sharp and sustained. In no jurisdiction is there clear evidence that growth rates are plateauing.

The total number of deaths is higher in the European jurisdictions than the North American jurisdictions, which is to be expected given that the former are more permissive.

To put the current number of deaths in context, in the Dutch speaking part of Flanders, physician-assisted deaths constituted 3.8% of deaths in 2007 and 6.3% of deaths (or 1 in 16) in 2013.23

A rate of over 6% of all deaths being carried out via physician-assisted deaths - and still rising - can be considered as nothing other than a major shift in practice and culture, particularly given that euthanasia was originally justified in Belgium as a last resort for the tiny minority of people for which palliation could not provide adequate care.

Table 2: Annual growth rate in cases of euthanasia and assisted suicide in selected jurisdictions

<table>
<thead>
<tr>
<th>Country</th>
<th>% Annual Growth in Total Deaths</th>
<th>CAGR24 in euthanasia / assisted suicide</th>
<th>No. of years over which CAGR calculated</th>
</tr>
</thead>
<tbody>
<tr>
<td>UNITED STATES</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oregon</td>
<td>0.525</td>
<td>13.226</td>
<td>17</td>
</tr>
<tr>
<td>Washington</td>
<td></td>
<td>21.627</td>
<td>6</td>
</tr>
<tr>
<td>EUROPE</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Belgium</td>
<td>0.328</td>
<td>19.629</td>
<td>12</td>
</tr>
<tr>
<td>The Netherlands</td>
<td>-0.330</td>
<td>13.131</td>
<td>7</td>
</tr>
<tr>
<td>Switzerland</td>
<td>0.332</td>
<td>19.233</td>
<td>16</td>
</tr>
<tr>
<td>AUSTRALIA</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Victoria</td>
<td>1.6134</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

24 Compound annual growth rate
27 Compound annual growth rate: 2009-2015
29 Compound annual growth rate: 2003-2015
30 http://stats.oecd.org/ - CAGR from 2000-2010 (latest data point)
31 Compound annual growth rate: 2008-2015
33 Compound annual growth rate: 1998-2015
Annual growth rates in the order of 13 – 20 per cent are extremely high. It is worth noting that the impact of cumulative growth rates of this magnitude over the medium term can be deceptive. What appears to be manageable at first can rapidly lead to very large numbers and, in the case of euthanasia and assisted suicide, a very high proportion of overall deaths.

For example, the total number of cases in Belgium has increased by 756% over 12 years and the number of cases in Oregon is 725% higher over the 17 years following legalisation.

Table 3 indicates what would happen in jurisdictions with legalised euthanasia and assisted suicide if currently observed growth rates continue. The final two columns indicate how many multiples of the base year frequency would occur after 10 and 20 years of sustained growth. These are not long periods of time.

For example, if Oregon’s observed annual growth rates were applied to a base number of 100 deaths, that number would grow to 350 after 10 years and 1,200 after 20 years. Higher growth rates, such as observed in Washington State, Belgium and Switzerland result in even more dramatic growth. Based on growth rates observed in Washington State, a base number of 100 cases in year 1 would grow to 700 ten years after commencement and 5,000 after 20 years.

It is not at all clear what kind of growth to expect in jurisdictions with legalised euthanasia and assisted suicide over the upcoming 10 or 20 years. However, we cannot rule out continued growth in the order of what we have already observed in jurisdictions such as Oregon and Switzerland.

Table 3: Annual growth rate in the cases of euthanasia and assisted suicide in selected jurisdictions

<table>
<thead>
<tr>
<th>Country</th>
<th>Observed annual growth rate by jurisdiction</th>
<th>Cumulative growth over 10 year period</th>
<th>Cumulative growth over 20 year period</th>
</tr>
</thead>
<tbody>
<tr>
<td>UNITED STATES</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oregon</td>
<td>13.2%35</td>
<td>3.5x</td>
<td>12x</td>
</tr>
<tr>
<td>Washington</td>
<td>21.6%36</td>
<td>7x</td>
<td>50x</td>
</tr>
<tr>
<td>EUROPE</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Belgium</td>
<td>19.6%37</td>
<td>6x</td>
<td>35x</td>
</tr>
<tr>
<td>The Netherlands</td>
<td>13.1%38</td>
<td>3.4x</td>
<td>12x</td>
</tr>
<tr>
<td>Switzerland</td>
<td>17.6%39</td>
<td>5.8x</td>
<td>34x</td>
</tr>
</tbody>
</table>

It is difficult to say, based on available data, why this is happening. Many jurisdictions with legalised euthanasia and assisted suicide do not provide for transparent reporting in relation to case details. The following chapter will outline five possible reasons for the rapid, sustained growth in the number of observed cases.

36 Compound annual growth rate: 2009-2015
37 Compound annual growth rate: 2003-2015
38 Compound annual growth rate: 2008-2015
CHAPTER 4 – REASONS FOR THE GROWING PREVALENCE OF EUTHANASIA AND ASSISTED DYING

There are many possible reasons why the number of cases of euthanasia and assisted suicide is rising so quickly and over such a long period of time. Understanding the causes of this growth is critical if we are to be confident that public policy objectives can be achieved through legalisation.

There is considerable evidence that this rapid and sustained rise in numbers is attributable in part to a growth in the scope of the practice, to a normalisation of the practices and to a failure of safeguards.

Box 1: Key public policy considerations

The Victorian Parliament should carefully evaluate evidence which demonstrates that:

- In jurisdictions where it is legal, access to euthanasia and assisted suicide is not being limited to those people experiencing unendurable physical pain, even where this was the original rationale for legalisation.
- Over time, the rapidly growing number of cases of euthanasia and assisted suicide reflects a disturbing normalisation of the practice, even though the original rationale for legalisation usually stressed that it was only intended to be used as a last resort in exceptional cases.
- Safeguards are systemically and routinely failing across a number of jurisdictions. Many of these safeguards are relied upon in the Majority Report without evidence that they will work in practice.

4.1 A greater awareness of rights unlikely to be a sustained driver of growth

Greater awareness of access to euthanasia or assisted dying could possibly have driven a growth in total cases in the months and even years following introduction of each scheme. As can be seen in Figures 1 to 5 however, high growth rates in total numbers have not abated in any jurisdiction, even where data exists for periods of almost two decades. In no jurisdiction is there any indication that growth rates are slowing. This suggests that sustained growth is due to factors other than community awareness.

4.2 An aging demographic contributes only a small share of overall growth

An aging demographic is often put forward as a possible reason why the number of cases is rising. Aging is not a major factor driving the growth in the number of cases of euthanasia or assisted dying in any of the jurisdictions for which data is available. As Table 2 shows, the total number of deaths over the period 2000-2010 grew at less than 1 per cent per annum in all relevant jurisdictions – and indeed it fell in the Netherlands. The change in the total number of deaths is less than one tenth of the total change in euthanasia and assisted dying cases.
4.3 Scope creep is occurring in at least some jurisdictions

There is a long-standing debate in ethics and jurisprudence as to the nature of the ethical difference (if there is one) between an act and an omission that lead to the same outcome. In the context of medical care, the debate centres on whether there is an ethical difference between a decision to end or withdraw treatment by a health practitioner and that same health practitioner actively taking part in ending someone’s life.

Regardless as to one’s views on whether there is a meaningful ethical difference between these two situations, the distinction between withdrawing treatment and undertaking a positive action is clearly defined. The logical and philosophical arguments on either side of debate are well developed and have been clearly articulated over centuries.

What is much less clear is the ethical difference between various types of action. For example, if one accepts that euthanasia or assisted suicide can be ethical in some situations, that raises subsequent questions such as whether it should be available only for people with a terminal condition or also for those suffering from a condition that is “unbearable”, albeit not terminal. Is physical pain necessary to trigger justifiable action or is psychological pain also sufficient? Is explicit consent required? If euthanasia or assisted suicide is justified only for people suffering from a terminal condition, what does “terminal” mean? Must death be expected within 3 months, 6 months, a year? Should an age limit be placed on consent? And so on.

The reason why the “slippery slope” argument has credibility in this context is that, once euthanasia or assisted suicide is accepted for a limited subset of situations, there is very little by way of logical or philosophical argument to stop the expansion of categories.

- If euthanasia or assisted suicide is permitted for those with a terminal condition in order to alleviate suffering – why not alleviate suffering of the same magnitude in those whose condition is not terminal?
- If consent is critical – then why limit availability to adults if “mature minors” can also give meaningful consent in other contexts such as joining the military.
- If advanced care directives can be used to justify the cessation of invasive and possibly painful medical procedures, why should advanced care directives not also be used to justify active euthanasia?

This is why the limitations included in euthanasia and assisted suicide regimes often come under immediate pressure as soon as these schemes are enacted. This takes two main forms. The first is attempts to explicitly expand the categories of situations in which euthanasia or assisted suicide is legal. The second is to reinterpret existing categories more broadly. As will be outlined below, reinterpretation has been undertaken in practice by both non-legislative and non-judicial bodies.

Often, legislation is passed with limitations that are designed to secure its passage through parliament. Where this “pragmatic” approach is adopted, these limits will almost inevitably be tested once the regime is in place.40

40 It is widely accepted that in Belgium, at least some of the restrictions contained in the initial legislation were included so as to secure passage of the Bill. Lemmens, Trudo, The Conflict Between Open-ended Access to Physician-Assisted Dying and The Protection of the
Some would argue that a slippery slope argument is flawed in that it doesn't acknowledge that subsequent changes might be justified on the basis of sound reasoning and policy formulation. That is true. There isn't anything wrong, per se, with legislation that has the potential to change.

The slippery slope argument is less concerned with the merits of each future potential change, and more with the inherent “momentum” in the system. The argument is that, once legalised, there will be a tendency for euthanasia and assisted suicide schemes to expand in scope since the logical and philosophical boundaries between different types of pro-active hastening of death are far weaker than the boundary between omissions and actions.

As will be argued below, the evidence bears this out in a number of jurisdictions.

4.3.1 Scope creep through legislative expansion of categories

Legislative changes have either been implemented or are under active consideration in a number of jurisdictions:

Belgium: euthanasia for children. The original euthanasia legislation in Belgium restricted access to people over the age of 18 years of age. On one reading, children were specifically excluded from the ambit of the legislation as “it was deemed so controversial that including it would have threatened approval of the Euthanasia Bill.” In 2014, an amendment was passed that allowed for euthanasia by “mature minors”.

Belgium: other changes under consideration by the legislature. In Belgium, a series of other legislative expansions of the euthanasia regime have been submitted to parliament, including:

- Requiring doctors to make referrals, even if it is against their conscience
- Permitting euthanasia for dementia patients
- The use of advanced care directives for people in an unconscious state

Canada: physician assisted dying for minors. While the Carter judgement by the Canadian Supreme Court only applied to competent adult persons, the Provincial-Territorial Advisory Group (PTAG) has already recommended allowing physician assisted dying for minors. Recommendation 17 of the PTAG Final Report is that:

Access to physician-assisted dying should not be impeded by the imposition of arbitrary age limits. Provinces and territories should recommend that the federal government make it clear in its changes to the Criminal Code that eligibility for physician-assisted dying is to be based on competence rather than age.

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42 European Institute of Bioethics, Euthanasia in Belgium: 10 years on, 2012, p4.

4.3.2 Non-legislative and informal extensions within an existing framework

Physician-assisted dying in Belgium is an example of informal slippage in the scope of the law. In his expert review for the Canadian Attorney-General, Professor Montero found that:

The Belgian legislator clearly intended to exclude physician-assisted suicide from the scope of the Act on euthanasia. This intention was criticized and debated extensively when the Act was being developed. A number of amendments were introduced to incorporate physician-assisted suicide into the scope of the Act, but they were all rejected to the Conseil d’Etat’s astonishment.

After the Act was passed, parliamentarians deemed it necessary to propose bills to amend the Act on euthanasia to include physician-assisted suicide performed under the same conditions as those applicable to euthanasia. To them, it is clear that physician-assisted suicide is not covered by the Act and therefore constitutes an illegal practice within the meaning of the Act as it is currently written.

Therefore, it is surprising that the Commission de controle regularly approves reported physician-assisted suicide cases and has been doing so since its first official report, stating that the practice “falls within the scope of the Act, as it is currently written ...”  

Lemmens argues that, in Belgium:

... the vague and open-ended nature of the terms ‘medically hopeless situation’ as well as a very subjective interpretation of the term ‘constant and unbearable physical or mental suffering’ have clearly opened the door to many instances of euthanasia that are controversial if not outright problematic.  

The Netherlands Organisation for Health Research and Development (ZonMw) published the second evaluation report on the functioning of the Termination of Life on Request and Assisted Suicide (Review Procedures) Act in 2012. The report finds that a more liberal approach is emerging in relation to requests from patients from some groups, such as those with mental illness or dementia.

This shift can be seen in medical opinion as well as in the policy position of the regional euthanasia review committees. The evaluation report states that:

... this development does not imply an expansion of the legal requirements: it should be seen as further conceptualisation of the meaning and scope of the requirements, that are formulated rather “openly” in the Act."  

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44 Montero, Professor Etienne, Affidavit to Lee Carter v The Attorney General of Canada et al, Supreme Court of Canada, No. 35591, pp52-53.


As is noted below, the number of instances of people with dementia and mental illness being euthanised in the Netherlands is rising rapidly. This could reflect regulatory creep.

On 7 January 2016, the *Netherlands Times* report that:

The Ministries of Public Health and Security and Justice updated their “guide” on euthanasia to give people suffering from severe dementia the option of euthanasia, even if they are unable to express the wish themselves. This is only possible if the patient wrote a declaration with his wish for euthanasia while he was still clearheaded, broadcaster NOS reports.47

The proposed physician-assisted dying regime in the Majority Report will establish an entity responsible for monitoring assisted dying within Victoria. Given the experiences of such entities in other jurisdictions, the risk that a supervisory body in Victoria will unilaterally extend the scope of any assisted dying regime through non-transparent interpretation of legislative provisions is a real risk and of great concern.

4.4 There is evidence of a "normalisation" of euthanasia and assisted suicide

Chapter 3 presents clear evidence of an upward and persistent trend in the number of cases of euthanasia and assisted suicide following legalisation. This section will refer to commentary that relates to a contemporaneous normalisation of the practice.

The term "normalisation" in this context is used to indicate that, once legalised, euthanasia and assisted suicide are increasingly taken for granted and seen to be unexceptional both within the medical profession and more broadly within society.

This is important given that the original rationale for euthanasia and assisted suicide in most jurisdictions, and in the Majority Report, is to deal with a set of circumstances that is extremely rare.

4.4.1 Belgium

Belgium currently has one of the most liberal euthanasia regimes in the world. Carine Brochier of the European Institute of Bioethics reported that one physician had told her:

You are right to say that many euthanasia cases are not [officially] declared. I myself have practiced euthanasia very many times, and I never declared them. It is too personal and a matter between the patient and his doctor. No need to publicise it.48

The open-ended regime in Belgium evokes "a culture of normalization of active life-ending interventions by physicians that may have long-term consequences which are hard to predict."49

In Belgium, the number of cases of euthanasia is rising rapidly. One would think that this would mean that a growing number of instances of euthanasia applications are falling into more difficult categories. But a recently published survey of physicians in Flanders found a rise in the proportion of requests granted from 56.3% in 2007 to 76.8% in 2013.50 This could indicate a growing level of comfort with physician-assisted dying amongst physicians.

Wim Distelmans, who was Chairman of the Federal Control and Evaluation Commission in Belgium, was interviewed by the New Yorker and stated that:

We at the commission are confronted more and more with patients who are tired of dealing with the sum of small ailments – they are what we call ‘tired of life’.


If you ask for euthanasia because you are alone, and you are alone because you don’t have family to take care of you, we cannot create family.51

In 2013, Wim Distelmans euthanised a 44 year old transgender man, Nathan Verhelst, because Verhelst was devastated by the outcome of his sex-change operation.52

Lemmens reports on a number of stridently pro-euthanasia physicians in Belgium who publicly argue for a need to widen current practices and who reject basic regulatory requirements such as reporting cases to regulators or consulting a second physician.53

A report by the European Institute of Bioethics warned that euthanasia has become normalised:

Initially legalized under very strict conditions, euthanasia has gradually become a very normal and even ordinary act to which patients are deemed “to have a right”. In the face of certain high profile cases, the evident relaxation of the very strict conditions has caused many reactions but also a total absence of any sanctions on the part of the Commission and a very conciliatory silence from the political establishment has given rise to a feeling of impunity on the part of some concerned medical practitioners, and to a feeling of powerlessness in those worried about where things are leading.54

4.4.2 The Netherlands

A recent article by a former review committee member in the Netherlands, Theo Boer, notes a shift in the type of patients who seek assisted dying:

... beginning in 2007, the numbers of assisted dying cases started going up by 15 percent each year. In 2014 the number of cases stood at 5,306, nearly three times the 2002 figure.

A shift has also taken place in the type of patients who seek assisted dying. Whereas in the first years the vast majority of patients—about 95 percent—were patients with a terminal disease who had their lives ended days or weeks before a natural death was expected, an increasing number of patients now seek assisted dying because of dementia, psychiatric illnesses, and accumulated age-related complaints.

Terminal cancer now accounts for fewer than 75 percent of the cases. Many of the remaining 25 percent could have lived for months, years, or even decades. In some reported cases, the suffering largely consists of being old, lonely, or bereaved.55

54 European Institute of Bioethics, Euthanasia in Belgium: 10 Years On, 2012, p7.
This normalisation manifests itself in many ways. In 2012, mobile euthanasia clinics began providing lethal injections to people free of charge at their homes in the Netherlands:\(^{56}\)

The launch of the so-called *Levenseinde*, or "Life End", house-call units – whose services are being offered to Dutch citizens free of charge – coincides with the opening of a clinic of the same name in The Hague, which will take patients with incurable illnesses as well as others who do not want to die at home.

The scheme is an initiative by the Dutch Association for a Voluntary End to Life (NVVE), a 130,000-member euthanasia organisation that is the biggest of its kind in the world.\(^{57}\)
4.5 A growing number of cases involving vulnerable people

One of key arguments used to support the legalisation of euthanasia and assisted dying is the need to respect peoples' autonomy. The growing number of cases involving very vulnerable people or people who may not have a capacity to provide meaningful consent raises serious questions about the validity of this rationale.

4.5.1 People who feel as though they are a burden

In 2006, Dr Harvey Chochinov undertook a review of the empirical literature related to the psychological, existential and spiritual aspects of patient deaths.\textsuperscript{58} Some key findings cited by Chochinov in relation to this issue were:

- A study of reports from family members of patients in Oregon and Washington State who expressed a wish for hastened death indicated that 59% of patients who did not discuss wanting assisted suicide and 94% of patients who did discuss wanting assisted suicide experienced distress due to feeling like a burden on others.\textsuperscript{59} The study found that: “Data from other studies and from the Oregon Health Division supports that fear of being a burden is common in patients who die by lethal prescription.”\textsuperscript{60}
- This is corroborated from survey data of caregivers in Japan.\textsuperscript{61}
- In a study from Northern Ireland, 103 doctors provided case history details of patients who had requested euthanasia. Being a “burden to others” was the second most commonly cited concern of patients, being raised in 54.9% of cases.\textsuperscript{62}
- A study of patients who killed themselves found that many were concerned about being a burden on others: “Multiple vulnerability factors were present simultaneously in all patients. However, the loss of, and the fear of losing, autonomy and their independence and of being a burden on others were the most relevant.”\textsuperscript{63}

A 2009 survey of 56 Oregonians who had either requested physician aid in dying (PAD) or contacted a PAD agency sought to explore the motivation for these requests. Respondents were asked to rate each of 29 possible reasons on a scale from 1 to 5 (1 being not very important through to 5 being very important). The most important reasons, with a median score of 5, included “wanting to

\textsuperscript{58} Chochinov, Harvey Max, "Dying, Dignity, and New Horizons in Palliative End-of-Life Care", CA Cancer Journal for Clinicians, 2006; 56:84-103.
\textsuperscript{60} Morita T, Sakaguchi Y, Hirai K, et al., “Desire for death and requests to hasten death of Japanese terminally ill cancer patients receiving specialized inpatient palliative care.”, Journal of Pain Symptom Management, 2004;27:44–52. This was a survey of 500 primary caregivers with a response rate of 290. It found that one of the major reasons for desiring death and for making a request to hasten death was being a burden on others.
\textsuperscript{62} Morita T, Sakaguchi Y, Hirai K, et al., “Desire for death and requests to hasten death of Japanese terminally ill cancer patients receiving specialized inpatient palliative care.”, Journal of Pain Symptom Management, 2004;27:44–52. This was a survey of 500 primary caregivers with a response rate of 290. It found that one of the major reasons for desiring death and for making a request to hasten death was being a burden on others.
control the circumstances of death”, “future poor quality of life”, “future pain” and “loss of independence”. Importantly, another reason that rated very high, with a median score of 4 was “perception of self as burden”. 64

This corroborated results from an earlier 2001 study of physician-assisted suicide in Oregon reported in the New England Journal of Medicine which found that 63% of patients who received a hastened death under the Dying with Dignity Act had expressed a strong sense of being burden and that “the frequency of concern about being a burden to others has increased.” 65

In the most recent report published by the Oregon Public Health Division, of the 132 deaths for which data was available, 48% listed being a burden on family, friends or caregivers as a concern. 66

Box 2: Case Study - Kate Cheney, Oregon

“Kate Cheney, an eighty-five-year-old widow, was diagnosed as terminally ill with stomach cancer. Kate wanted the option of assisted suicide in case she was in pain or if the indignities of losing control of her body functions became unbearable. Her daughter Erika, a retired nurse who had come from Arizona to care for her mother, went with Kate when she made her request for assisted suicide to her physician at Kaiser Permanente. Erika described the physician as ‘dismissive’ and requested and received a referral to another Kaiser physician. Kate’s second doctor arrange for a psychiatric consultation, a standard procedure at Kaiser. Although the psychiatrist who visited Kate at her home declined to be interviewed, the family released his report to the Oregonian’s reporter. The psychiatrist found that the patient did ‘not seem to be explicitly pushing for assisted suicide’ and lacked ‘the very high level of capacity to weigh options about it.’ Although the patient seemed to accept the assessment, the psychiatrist noted that the daughter became very angry.

Kaiser then suggested that the family obtain a second assessment from an outside consultant. The psychologist consulted noted that Kate had some memory defects and that her ‘choices [might have been] influenced by her family’s wishes, and that her daughter, Erika, [might have been] somewhat coercive’ but felt that Kate had the ability to make her own decision. A Kaiser administrator saw Kate and decided that she was competent and was making the decision on her own. Kate received the lethal drugs, which were put under Erika’s care.

As time went by and Kate ate poorly and became somewhat weaker, Erika and her husband needed a respite and sent Kate to a nursing home for a week. Kate ate well there, but when Erika visited Kate always asked when she would be going home. On the day she returned from the nursing home she told Erika and her husband that something had to be done, given her declining health. She had considered going permanently into a nursing home but decided against it. She told them she wanted to use the pills and asked for their help. ‘When would you like to do this?’ her son-in-law asked. ‘Now’, Kate replied. Grandchildren were contacted, those who lived nearby came over, goodbyes were said, and within a short time, with her family beside her, Kate took the pills and died.” 67

65 Sullivan AD, Hedberg K, Hopkins D., “Legalized physician-assisted suicide in Oregon, 1998-2000”, New England Journal of Medicine, 2001;344:605-607. The percentage of patients with concerns about being a burden on family, friends or other caregivers rose from 2/16 (12%) to 17 /27 (63%).
In the case of Kate Cheney, one can easily see how an elderly person could feel as though they were a burden on their family. The pressure on Kate's family was immense and her daughter and son-in-law were in desperate need of respite. However, it is also true that, as Foley and Hendin observe, "one can readily see how in the best of circumstances frail elderly patients can feel coerced to die."\(^{68}\)

### 4.5.2 Depression

Many people experiencing terminal diseases also suffer from depression. This is completely unsurprising. The prevalence of depression in people with terminal conditions raises a significant question as to how that depression interacts with decision-making. It goes without saying that experiencing depression doesn't necessarily make one unable to make sound decisions. But depression can impact on decision-making in a way that raises questions about the effectiveness of safeguards in relation to irreversible choices.

Depression is common amongst those seeking euthanasia and assisted suicide. In a study of 138 cancer patients with an estimated life expectancy of 3 months or less, a strong connection was found between depression and a request for euthanasia. Specifically, a patient was 4.1 times more likely to request euthanasia if depressed than if not depressed.\(^{69}\)

A study of suicide victims who had suffered from cancer found that depressive syndromes were present in 80 per cent of cases. "Only a small minority of cancer suicides seem to occur in the absence of mental disorders."\(^{70}\)

In a study of 44 terminally ill patients by Brown et al, 34 had never wished death to come early and ten were either suicidal (3) or desired an early death (7). All ten of the patients who desired an early death were found to be suffering from clinical depressive illness.\(^{71}\)

Ganzini et al, in a broad ranging review of instances of assisted dying in Oregon, found that twenty percent of the patients had symptoms of depression.\(^{72}\)

These studies raise questions about what safeguards need to be in place in legalised euthanasia or assisted suicide regimes to protect people from making decisions that might be affected in the short term by a depressive mood that might be manageable or treatable with additional support.

Some argue that euthanasia and assisted suicide are suitable in cases of treatment-resistant depression. Even in these cases, treatment options are often available. A study of the longer-term outcomes for patients with confirmed treatment-resistant depression examined the outcomes between 8 to 34 months following treatment in a specialist in-patient centre. The study included 118 participants. The key result was that the majority of participants (60.2%) were in full remission.

\(^{68}\) Ibid, p157.

\(^{69}\) Van der Lee, Marije, Johanna van der Bom, Nikkie B. Swarte, A. Peter Heintz, Alexander de Graeff and Jan van den Bout, "Euthanasia and Depression: A Prospective Cohort Study Among Terminally Ill Cancer Patients", *Journal of Clinical Oncology*, Vol 23, Number 27, 2005, pp6608-6609.


following specialist care and that nearly half of the total sample (48.3%) maintained full remission for at least 6 months. 73

While this does not mean that all cases of treatment-resistant depression can be fully treated, it does suggest that great care needs to be taken where a person is suffering from depression and is seeking either euthanasia or assisted dying. Given the range of treatments currently available and the ever improving prospects for treatment over time, the risk of allowing someone to choose death when realistic treatment options are available is a real one.

A number of studies have shown that physicians find it difficult to diagnose depression in patients with terminal conditions, let alone to determine whether that depression is impairing judgement. 74 Foley and Hendin argue that the Ganzini et al study cited above of physicians administering lethal injections in Oregon suggests potential difficulties in diagnosis. In that study of 143 cases, 20 per cent were identified as having depression which Foley and Hendin argue is materially well below the more commonly established rate of approximately 60 per cent or more. 75 This suggests that at, in at least some cases in the survey, depression may have been undiagnosed.

In 2015, The New Yorker examined the high profile case of Godelieva De Troyer in considerable detail. 76 This is a complex case from Belgium involving a woman who had suffered from severe depression over a long period of time. During a period of alienation from her adult children, she was euthanised without their knowledge. This case has achieved a high degree of public attention both within Belgium and internationally as it highlights the particularly complex circumstances that can arise when euthanasia is carried out on a person with a mental health condition.

One of the main reasons that the case has generated public debate is the campaign that was subsequently waged by her son. He has challenged the appropriateness of a person being euthanised without family members being consulted. He argues that is was particularly inappropriate in Godelieva’s case given that her depression arose, in part, as a result of estrangement from her family. A reconciliation might have been a key element in her recovery. Issues relating to consultation with family members in cases of mental health care decisions arise more broadly than in cases of euthanasia. But these issues are particularly pronounced in euthanasia given the irreversibility of the decision.

A second issue relating to this case that has caused public controversy is a review of Godelieva’s medical file that was undertaken by Dr Georges Casteur, a former president of the provincial council of the doctors of West Flanders. His review found that Godelieva had “struggled to find three doctors who would say that she had an incurable illness, as the law required. One psychiatrist wrote that her desire for euthanasia was ‘not mature’, because she has ‘ups and downs’”. According to the reviewing psychiatrist, a second doctor concluded that she could still be helped and that “when

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Godelieva discussed her grandchildren she became emotional and expressed doubts about her decision to die.\textsuperscript{77}

The purpose of raising this case is not to dwell on the particulars of Godelieva’s situation, but rather to highlight the difficulties in diagnosis and treatment in cases involving depression and mental health issues more generally. Jurisdictions such as Belgium and the Netherlands have more stringent safeguards in place where there is a mental health condition, but the effectiveness of these safeguards in practice is not proven.

4.5.3 Mental illness

This paper will not examine the merits of whether a scheme should permit euthanasia or assisted suicide where a patient is suffering from a mental illness. At the very least, it is worth noting that in such situations additional care should be taken as diagnosis will often be challenging and it will usually be more difficult to clearly establish consent. The growing prevalence of euthanasia and assisting dying for people with mental illness in at least some jurisdictions raises concerns about the effectiveness of safeguards.

The Belgian legislation allows those experiencing mental suffering to access physician assisted dying but includes additional safeguards for people in this category. This includes a detailed consultation with a third physician and a mandatory one month waiting period between the request and the life-ending act.

The proportion of euthanasia deaths involving neuropsychiatric disorders has increased sharply in Belgium over the past decade, from 1.2\% of cases in 2004/05 to 2.8\% in 2010/11 (58 cases) and 3.7\% of cases in 2013/14 (67 cases).\textsuperscript{78} Table 4 contains the number of instances of euthanasia in Belgium over recent years as reported to the Belgian legislature.

Table 4: Number of cases of euthanasia for neuropsychiatric conditions in Belgium\textsuperscript{79}

<table>
<thead>
<tr>
<th>Source</th>
<th>Years covered by report</th>
<th>Number of cases of neuropsychiatric conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Second report</td>
<td>2004 and 2005</td>
<td>9\textsuperscript{80}</td>
</tr>
<tr>
<td>Third report</td>
<td>2006 and 2007</td>
<td>13\textsuperscript{81}</td>
</tr>
<tr>
<td>Fourth report</td>
<td>2008 and 2009</td>
<td>62</td>
</tr>
<tr>
<td>Fifth report</td>
<td>2010 and 2011</td>
<td>105</td>
</tr>
</tbody>
</table>

\textsuperscript{77} Aviv, Rachel, “The Death Treatment”, June 22 2015, The New Yorker, \url{http://www.newyorker.com/magazine/2015/06/22/the-death-treatment}


\textsuperscript{79} Montero, Professor Etienne, Affidavit to Lee Carter v The Attorney General of Canada et al, Supreme Court of Canada, No. 35591, p51.

\textsuperscript{80} This included three cases of Alzheimer’s disease, one case of Huntington’s disease, one case of Creutzfeldt-Jacob disease and four cases of unyielding depression.

\textsuperscript{81} Nine cases of neuropsychiatric conditions (dementia or depression) in addition to four for people with neuropsychiatric conditions for whom death was not expected in the short term.
In the Netherlands, recent data from reports of the Regional Euthanasia Review Committees points to a growing number of cases of euthanasia in cases of mental illness and dementia. Table 5 contains the number of cases of mental illness and dementia over the period 2012-2015.

While the number of cases in each category is coming off a low base, the rapid CAGR rate suggests that number of cases in each category could be very large in the not too distant future. There is no reason to think that growth rates in either category will taper off given what we observe in growth rates in the overall number of cases both in the Netherlands and other major jurisdictions.

Table 5: Number of cases of euthanasia for mental illness or dementia in Netherlands

<table>
<thead>
<tr>
<th>Year</th>
<th>Mental Illness (Cases)</th>
<th>Dementia (Cases)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2012</td>
<td>14</td>
<td>42</td>
</tr>
<tr>
<td>2013</td>
<td>42</td>
<td>97</td>
</tr>
<tr>
<td>2014</td>
<td>41</td>
<td>81</td>
</tr>
<tr>
<td>2015</td>
<td>56</td>
<td>109</td>
</tr>
</tbody>
</table>

Growth rate: 59% 37%

As is noted above, when the Belgian Act on euthanasia was being debated and adopted, it was stated repeatedly that patients with psychiatric disorders, dementia or depression would be excluded from the Act. However, as in the Netherlands, there has been a rapid growth in this category over recent years.

In 2007, the Netwerk Depressie Vlaanderen, a Flemish association that provides support to people suffering from depression, stated that: “The door to euthanasia is open for thousands of depressed and suicidal people to kill themselves legally.”

Dealing with requests for euthanasia or assisted suicide from people with mental illness is particularly challenging for medical practitioners. Kim et al undertook a detailed review of 66 cases of euthanasia and assisted suicide involving psychiatric conditions in the Netherlands between 2011 and 2014. Two senior psychiatrists reviewed reports provided by the Dutch regional euthanasia review committees up to 1 June 2015. They found that:

- Most patients had chronic, severe conditions with histories of attempted suicides and psychiatric hospitalisations.
- A range of conditions were present including: depression; personality disorders; psychotic, post-traumatic stress or anxiety; neurocognitive and eating disorders; as well as prolonged grief and autism.
- 27% (n=18) of patients received euthanasia or assisted suicide from physicians new to them.

82 See Dutch Regional Euthanasia Review Committee Annual Reports.
83 Montero, Professor Etienne, Affidavit to Lee Carter v The Attorney General of Canada et al, Supreme Court of Canada, No. 35591, p51.
84 Statement made in the article “Quatre cas pour depression majeure irreductible”, La Libre Belgique, 2 February 2007. See translation in Montero, Professor Etienne, Affidavit to Lee Carter v The Attorney General of Canada et al, Supreme Court of Canada, No. 35591, p51.
• Of the 18 physicians that were new to patients, 14 of these were from the End-of-Life Clinic, a mobile euthanasia clinic. In 2012, the NVVE (Dutch Association for a Voluntary End to Life) created the End-of-life Clinic and started making “house calls” via mobile euthanasia units that respond to the wishes of patients whose regular doctors have refused to carry out their wishes.

Kim et al conclude that the “granting of their EAS [euthanasia or assisted suicide] requests appears to involve considerable physician judgment, usually involving multiple physicians who do not always agree (sometimes without independent psychiatric input)”.

The difficulty of evaluating suicidal patients with serious medical illness is explored by Hamilton et al. They argue that there are competing paradigms: the traditional model; and the assisted suicide competency model.

In the Harvard Medical School Guide to Suicide Assessment and Intervention, Hendin argues that medically ill suicidal patients are no different from other suicidal individuals. “Although physical illness may be a precipitating cause of despair, these patients usually suffer from treatable depression and are always ambivalent about their desire for death.”

Hamilton et al argue that “when it comes to treatment, the approach in this population emphasizes an effort to ‘understand and relieve the desperation that underlies the request for assisted suicide.’ To do so, the clinician must resist assuming the role of ‘gatekeeper’, someone who would focus on issues of competence alone.”

This approach, they argue, can be contrasted with the assisted suicide competency model. One example of the alternative approach is contained in Oregon’s assisted suicide guidebook which states that:

The mental health consultation as outlined in the Oregon Act, is a form of a capacity or competence evaluation, specifically focused on capacity to make the decision to hasten death by self-administering a lethal dose of medication.

In the assisted suicide competency model, there is no obligation to treat depression or mental illness where it is found.

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The differences between the approach to dealing with patients suffering from depression and mental illness in Oregon assisted suicide consultations (and in similar jurisdictions) as compared to more traditional treatment clinical models is telling. It is potentially problematic in an area where there is still considerable disagreement between experts as to the relationship between depression, mental health issues and competence.

4.5.4 Cases in which explicit consent is not given

One of the difficulties of drawing conclusions in relation to life-ending acts without explicit patient request is the lack of clear data. In a large-scale death certificate survey, Chambaere et al find that “life-ending acts without explicit request are significantly different from those provided in euthanasia and similar to those provided in standard palliative care.” 93 This suggests that some of the worst fears that have been expressed in this area are based more on poor reporting than systemic abuse.

However, they also argue that legalisation of euthanasia or physician-assisted dying did not put an end to the practice of non-voluntary or involuntary termination of life (as argued elsewhere in this paper) and that: “we recommend a more nuanced view of life-ending acts without explicit patient request in the debate on physician-assisted dying.” 94

Some advocates claim that legalisation of either euthanasia or assisted dying will bring practices such as euthanasia without consent into the open. Seale examines end-of-life decisions in the UK and concludes that the rate of this type of end-of-life decision-making is sufficiently low that: “in the UK, this argument cannot be made.” 95

In contrast, it appears that this type of euthanasia has not disappeared from countries which have legalised euthanasia or assisted suicide, with relatively high rates persisting in Belgium. A study in the Lancet by Van der Heide et al undertook surveys in six European countries to contrast end-of-life decision making. It found that ending of a patient’s life without request occurred at rates ranging from 0.06% (Italy) through to 1.50% (Belgium). 96 Two countries with the highest rates of this type of end-of-life (Belgium at 1.5% and the Netherlands with 0.60%) allowed the practice of euthanasia and assisted dying.

Moreover, as will be argued below, it is clear that wherever the legal line is drawn, activity occurs in the grey area around that line that is difficult to monitor.

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4.5.5 Patients receiving inadequate pain relief or palliative care

Many situations in which a person requests a hastening of death involve pain management or palliative care that fall short of best practice. Before legislating for euthanasia or assisted suicide, it would be appropriate to provide greater support to health care providers dealing with patients at the end of life.

Cases involving inadequate pain management

The Committee received considerable expert evidence that best practice medical treatment and palliative care can now effectively manage pain in almost all cases. Occasionally, best practice is not followed. The best response to the occasional failure to maintain best practice is to provide appropriate resourcing to the health care system – including training for health practitioners - and to ensure that processes to monitor the quality of end-of-life care are adequate.

Dr Natasha Michael, Director of Palliative Care at Cabrini Health argued that, many of the instances of poor end-of-life care that are used to justify euthanasia or assisted suicide reflect practitioners who:

... practice outside of the scope of their training and experience and neither have the knowledge, experience or skill to assist those who suffer. The ongoing under-resourcing of psychosocial support to patients and family needs urgent attention in palliative care and in end-of-life care. The association between the expression of a desire for death and depression, anxiety and demoralisation is well-documented in international research. However, there remains poor availability of key staff in clinical psychology, liaison psychiatry and social work services in our populations.97

Cases involving inadequate palliative care: Oregon

In a study published in the New England Journal of Medicine, Ganzini, Nelson, Schmidt, et al. reported the results of an extensive survey of physicians eligible to prescribe lethal medications under the Oregon Death with Dignity Act. Of the 2,649 physicians responding (65% response rate), 144 had received a total of 221 requests for prescriptions of lethal medications. Complete information was received for 143 patients, including the final outcome for those patients.98

In the case of 68 patients, the treating physician implemented at least one substantive intervention, including: control of pain or other symptoms; referral to a hospice; a mental health, social work, chaplaincy or palliative care consultation; or a trial of antidepressant medication. Of the patients who experienced a substantive intervention, 31 out of 67 (46%) changed their minds about wanting a prescription for lethal medication as opposed to 11 out of 73 (15%) for whom no substantive

97 Dr Natasha Michael, Director of Palliative Care, Cabrini Health, Transcript of evidence, 16 September 2016, pp7-8.
intervention was given. This material difference in outcomes indicates how dangerous it can be if any patients fail to access all best practice treatment interventions when they are in a vulnerable situation and considering suicide or euthanasia.

The 2016 Report of the Oregon Health Division indicates that 28.7% expressed inadequate pain control or a concern about it as a reason for wanting assisted suicide. This was the second least commonly cited reason in 2015.

It would be useful to know, of the 28.7%, what proportion were experiencing unendurable pain at the time the lethal dose was prescribed and administered and what proportion were suffering from an anticipation of future pain. It would also be useful to know, of those patients experiencing unendurable pain at the time the dosage was administered, how many were not benefiting from best practice pain relief of palliative care.

Advocates of euthanasia and assisted dying often point to the potential for legalised regimes to provide people with comfort by providing them with the assurance that they will be able to opt for a painless death if they wish to. Yet how many people under the assisted dying regime in Oregon (and similar regimes) secure access to lethal drugs out of a fear of future unbearable pain only to use those drugs before they have explored best practice palliative care options? Given current reporting arrangements, it is impossible to know with confidence.

Cases involving inadequate palliative care: the Northern Territory

The legalisation of euthanasia in the Northern Territory was accompanied by a number of safeguards that advocates of euthanasia and assisted suicide claim can protect vulnerable people in practice, including that:

- Under the legislation, an opinion from a second medical practitioner was required to verify the existence and terminal nature of the patient’s illness. The second medical practitioner was required to have “special expertise in the illness [being experienced by the patient] and qualifications in a medical specialty recognised by fellowship in a specialist college in Australia.”
- If the first doctor did not have special qualifications in palliative care, a third doctor with such expertise and qualifications was required to give information to the patient on the availability of palliative care.
- A psychiatrist was required to examine the patient to certify that he or she did not have “a treatable clinical depression.”

The outcomes of this brief episode of legalising euthanasia are highly instructive as it is the only example of legalisation in Australia to date.

102 Ibid, p194.
103 Ibid, p194.
The experiences of people whose lives ended under the Northern Territory regime casts doubt on the effectiveness of the types of safeguards proposed in the Majority Report. During the 9 months that euthanasia was legal in the Northern Territory (July 1996 – March 1997), seven people made formal use of the Rights of the Terminally Ill (ROTI) Act, four of whom died. All seven patients had cancer, most at an advanced stage.

In a paper by Kissane, Street and Nitschke, it was found that three of the seven patients were socially isolated and depressive symptoms were present in four of the seven. Failure of pain management did not appear to be the motivation for the patients. Of the seven patients, four had controlled pain and the remaining three did not have prominent pain. This is in contrast to the way this issue is often framed by advocates of euthanasia and assisted dying.

The requirement that there be a consensus as to whether the patient was terminal was problematic. There was a consensus that the patient was terminal in four cases, a lack of consensus in two cases and the patient was not terminal in one case. In one instance, an oncologist gave a prognosis of 9 months and a dermatologist and local oncologist both judged that the patient was not terminal.

The ROTI Act contained a requirement that a psychiatrist confirm that the patient was not suffering from a treatable clinical depression. "Confirmation was not easy since patients perceived such a mandatory assessment as a hurdle to overcome." To what extent was the psychiatrist trusted with important data and able to build an appropriate alliance that permitted a genuine understanding of a patient’s plight? In case 1, there was important background detail about the death of one child and alienation from another, which was withheld during the psychiatric assessment. These experiences may have placed the patient in a lonely, grieving, demoralised position: an unrecognised depression may have led to suicide.

Four of the seven cases had symptoms of depression, including reduced reactivity, lowered mood, hopelessness, and suicidal thoughts. Case 4 was receiving treatment for depression, but no consideration was given to the efficacy of dose, change of medication, or psychotherapeutic management. PN judged this patient as unlikely to respond to further treatment. Nonetheless, continued psychiatric care appeared warranted—a psychiatrist can have an active therapeutic role in ameliorating suffering rather than being used only as a gatekeeper to euthanasia. Ganzini and colleagues showed that only 6% of psychiatrists in Oregon, USA, thought that they could be a competent gatekeeper after a single assessment of a patient.

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105 This paper, which appeared in the Lancet, was based upon, inter alia, 18 hours of interviews with Nitschke by Professor David Kissane, a psychiatrist and professor of palliative care and Annette Street, a medical sociologist. In addition, the authors had access to documents from the coroner’s court, public texts created by the patients and other public commentary (eg in the media).
106 Kissane, David, Annette Street and Philip Nitschke, "Seven Deaths in Darwin: case studies under the Rights of the Terminally Ill Act, Northern Territory, Australia", The Lancet, Vol. 352, October 3 1998, p1098. Two had depressive symptoms (one of whom was on antidepressants), one had a history of depression and was experiencing suicidal thoughts, and another was suicidal (this person ultimately died as a result of suicide).
107 Ibid, p1098.
110 Ibid, pp1101.
In a follow up article, Professor Kissane examined each of the seven cases briefly summarised above. He found that demoralisation, which could at least partly be managed, played a significant role:

Review of these patients' stories highlighted for me the importance of demoralization as a significant mental state influencing the choices these patients made. It is likely that the mental state of demoralization influenced their judgement, narrowing their perspective about available options and choices. Furthermore, demoralized patients may not make a truly informed decision in giving medical consent.

The prognostic language within oncology that designates 'there is no cure' is one potential cause of demoralization in these patients, a cause that be avoided by more sensitive medical communication with those who are seriously ill. 111

Kissane also found that other perceptions were common in patients, most of which were manageable through effective medical and palliative care:

... this Australian cohort considered concern about loss of dignity, becoming dependent on others, and potentially being a burden as prominent reasons for the request for euthanasia. ...

Research has repeatedly shown how quality of life is appraised differently by patient, caregiver and clinician. A patient with cancer can adjust to the experience of gradual frailty over time, so long as adequate reassurance is given about the thoroughness of care long the way. 112

The experience of euthanasia in the Northern Territory demonstrates that a regime with what are typically considered to be strong safeguards can fail vulnerable people. In the Northern Territory, people who experienced social isolation, isolation from family, depression and demoralisation were not given the best possible treatment.

4.5.6 Summary of evidence in relation to vulnerable people

There is considerable evidence that many vulnerable people will be put into a difficult position if euthanasia or assisted suicide are choices that become a standard part of the set of treatment options available at the end-of-life.

The House of Lords Select Committee on Ethics came to a similar overall conclusion:

... it would be next to impossible to ensure that all acts of euthanasia were truly voluntary and that any liberalisation of the law was not abused.

vulnerable people—the elderly, lonely, sick or distressed—would feel pressure, whether real or imagined, to request early death113

112 Ibid, p204.
4.6 The widespread failure of safeguards

The Majority Report asserts that the evidence is “clear” that safeguards work in jurisdictions with legalised euthanasia and assisted suicide. A balanced reading of the evidence would lead one to conclude that such an unequivocal statement is not true.

4.6.1 Systemic non-reporting

A requirement to report instances of euthanasia and assisted suicide is seen by some advocates as a key strength of a legalised regime. They argue that it will bring opaque practices more into the open. The experience in Belgium suggests otherwise.

In Belgium, mandatory notification of euthanasia to the Federal Control and Evaluation Commission is a cornerstone of the regulatory arrangements. However, recent reports suggest that around half of all euthanasia cases are not reported. In one study, physicians who did not report cases of euthanasia provided the following responses:

- Did not perceive their action to be euthanasia: 77%
- Too much of an administrative burden: 18%
- That legal due care requirements had not been met: 12%
- That euthanasia is a private matter: 9%
- Possible legal consequences: 2%

The high rate of non-reporting and the reasons given are a matter of serious concern and raise doubts about the effectiveness of official oversight.

While the rate of non-reporting is falling in the Netherlands, it appears to be of a similar magnitude to that found in Belgium.

A low rate of reporting could potentially lead to serious consequences. A cross sectional analysis of reported and unreported euthanasia cases provided evidence that unreported cases were generally dealt with less carefully than reported cases. Specifically, in unreported cases: a written request for euthanasia was more likely to be absent; independent physicians and caregivers specialising in palliative care were consulted less often; the life ending act was more likely to be performed with opioids or sedatives; and the drugs were more often administered by a nurse, not a physician as required.


115 Multiple answers are possible, which is why the various categories sum to more than 100 per cent. Smets T., J. Bilsen, J. Cohen, M.L. Rurup, F. Mortier, L. Deliens, “Reporting of euthanasia in medical practice in Flanders, Belgium: Cross sectional analysis of reported and unreported cases”, BMJ, 341 (2010), p. c5174.


The FCEC itself has stated that it is "not capable of assessing the proportion of declared cases of euthanasia compared with the number of real cases which have actually taken place."\(^{118}\)

There are numerous high-profile and well-documented cases of actions beyond the scope of the law in Belgium that are neither being reported nor referred to prosecutor. One recent example was a public interview given by Dr Mark Cosyns in which he concedes that he routinely fails to report and in which he details a recent case of euthanasia that was of questionable compliance.\(^{119}\) As noted above, Carine Brochier reports of a physician who stated that he routinely fails to declare instances of euthanasia.\(^{120}\)

In Oregon, low rates of reporting are a consequence of a regulatory regime that does not require sufficient scrutiny. According to Foley and Hendin, the Oregon Health Division has interpreted its mandate narrowly:

> OHD limits its yearly reports to general epidemiological data and collects limited information from physicians who have prescribed lethal medication. Physicians who declined to prescribe the lethal medication, as well as nurses and social workers who cared for patients, are not interviewed. ... There is no provision for an independent evaluator or researcher to study whatever data are available. This OHD process has presented a full and open discussion.\(^{121}\)

### 4.6.2 Confusion amongst physicians

A survey conducted among physicians in Belgium seven years after the legalisation of euthanasia demonstrated that there was little consensus among physicians as to how to label hypothetical end-of-life decisions or what reporting obligations attach to such decisions. The study was based on a survey of 3,006 physicians who had graduated in their area of specialty at least 12 months prior to the survey.

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\(^{119}\) Lemmens, Trudo, *The Conflict Between Open-ended Access to Physician-Assisted Dying and The Protection of the Vulnerable: Lessons from Belgium’s Euthanasia Regime in the Post-Carter Era*, (forthcoming), Catherine Regis, Lara Ighoury & Robert Kouri, eds., *Key Conflicts in Health Law*, (Cowensville: Yvon Blais, 2016), p.25 - See for example: an interview in *De Standaard* with Dr. Mark Cosyns, a palliative care specialist who has performed many PADs. When he is asked "Dr. Cosyns, you are still not reporting euthanasia to the Commission?" his answer is: "No, not when they are our own patients. I do everything on the basis of the Act on Patient’s Rights [another piece of legislation]. We also don’t have to justify each time we do an operation why this doesn’t constitute assault." ["Nee, niet als het om onze eigen patiënten gaat. Ik doe alles op basis van de wet op de patiënterrechten. We moeten toch ook niet voor elke operatie verantwoorden dat we geen opzettelijke slagen en verwondingen toebrengen"] Veerle BEEL and Lieven SIOEN, “In de VS zat ik al lang achter de tralies. En jij samen met mij” [In de VS I’d already be in prison a long time ago. And you with me], *De Standaard*, 21 December 2013, online: <http://www.standaard.be/cnt/dmf20131221_00896009>. In another interview, he admits, among other things, to have provided PAD to an elderly couple. “One of them”—to use his own words—“because of medical, terminal reasons and the other because of the psychological suffering of being old and having to continue living on her own after having lived nearly in symbiosis” [my translation of: "De ene om medische, terminale redenen en de ander omwille van het psychisch lijden van oud zijn en alleen verder te moeten na bijna in symbionse geleefd te hebben" quoted in Fred VERBAKEL, “Niet is zo natuurlijk als sterven”, (2009) 2:4 Relevant 5, 6, available online: <https://www.nvve.nl/files/8513/8753/089S/Relevant_2009-2.pdf>].


The report found considerable confusion in relation to classification of clear hypothetical situations:

... there is a lack of agreement among physicians in Belgium about the classification of euthanasia and other ELDs, and about which cases must be reported as euthanasia to the Federal Review Committee. Seven years after implementation of the euthanasia law in Belgium, 2 out of 10 physicians, likely to be involved in the care of dying patients, did not label a hypothetical case in which a physician ends the life a patient at that patient’s explicit request using neuromuscular relaxants as 'euthanasia'. Three out of 10 physicians did not know that the case had to be reported to the Federal Review Committee.

Most physicians labelled the euthanasia case in which the physician ends the life of a patient at that patient’s explicit request using morphine (case 3) as ‘intensification of pain and symptom treatment’ (39%) or as ‘palliative/terminal sedation’ (37%); only 21% of physicians labelled this case as ‘euthanasia’.122

Even among those physicians who labelled instances of euthanasia correctly, a considerable number did not know that they had an obligation to report such cases. This is likely to be a contributory factor to the low overall rates of reporting observed in Belgium.

4.6.3 The difficulty of deciding whether a case is terminal

Some euthanasia and assisted dying regimes limit eligibility to terminal cases. This might involve specifying a time-limit (e.g. a patient’s prognosis is that they are likely to die within, say, 6 months) while others are more open-ended. In practice, deciding whether a patient is “terminal” is extremely difficult. Physicians find it difficult to put specific time estimates on many patients’ conditions and, unsurprisingly, often disagree with each other.

In Oregon, a patient is only eligible for assisted suicide if they are terminally ill, with less than six months to live. When surveyed, over 50 per cent of Oregon physicians indicated that they were not confident that they could make such a prediction.123 It is unclear at present how this uncertainty is communicated to patients seeking assisted suicide.

Compounding this difficulty in Oregon is the fact that the proportion of cases involving advanced cancer is trending down. There is an increasing tendency for patients seeking lethal prescriptions to have conditions with a less predictable future trajectory such as ALS (16% in 2014), chronic lower respiratory disease, Multiple Sclerosis, Parkinson's disease, diabetes and heart disease.124 Of the 105 patients who took lethal drugs in 2014, 11 had been diagnosed as having less than six months to live in 2012 or 2013.125

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4.6.4 Doctor shopping

Foley and Hendin explore a case study in Oregon involving a woman in her mid-eighties with metastatic breast cancer who was in a hospice program. Given the lack of transparency surrounding assisted dying in Oregon, the information that they gathered included: reports from the staff of Compassion in Dying (the organisation that facilitated the assisted death); a news conference following the death; and information from anonymous interviews by the physician who prescribed the medication. According to these sources, the patient's own physician was not willing to assist in her suicide (for reasons not specified), a second physician refused on the basis that the patient was depressed and it was only when Compassion in Dying was contacted that a doctor was found who was willing to prescribe a lethal medication. The patient was seen by a psychiatrist - but only once.\(^{126}\)

While legally proscribed procedures appear to have been followed, the case raises a number of questions. First, the views of the patient's own doctor and the physician who believed that she was suffering from depression were essentially ignored. All that was required was for the patient to find another doctor willing to undertake the procedure. Any regime that permits such doctor shopping is prone to abuse. This kind of outcome is possible in a number of the euthanasia and assisted dying regimes currently in place.

This is similar to what occurred in the case of Godelieva De Troyer and is reflected in a study in Belgium which found that in 23 per cent of cases, euthanasia was performed despite a divergence of opinions.\(^{127}\)

4.6.5 Nurses administering medicine

Recent studies provide evidence that in Belgium nurses commonly administer fatal doses in contravention of the law. One recent study surveyed 6,000 nurses in Flanders. Of 1,265 completed surveys, 128 nurses reported that the last patient in their care for whom a life-shortening end-of-life decision was made received euthanasia and 120 reported that the patient received life-ending drugs without his or her explicit request. Of the nurses involved in euthanasia, 64% were involved in the decision-making process. The drugs were administered by the nurse in 12% of cases, mostly without the physician co-administering (12/14 = 86%).\(^{128}\) In using life-ending drugs without explicit request, 48% of nurses helped prepare drugs, 56% were present during administering and in 45 instances, the nurse administered the drugs, mostly without the physician co-administering (82%) but under physician orders (98%).\(^{129}\)


Another study focused on situations in which patients were euthanised without having made an explicit request to die. This study found that, in almost half of the cases (6 of 13), the final drug was administered by a nurse.\textsuperscript{130}

The administration of drugs by nurses is not a major concern per se. Nurses are often the key caregivers at the end of life with the most intimate knowledge of a patient’s preferences and needs. What is relevant in the current context is the fact that this practice is prevalent in a situation that is highly sensitive (the administration of fatal doses) and where the law clearly states that it should not occur. Clearly, in regimes where euthanasia and assisted suicide are legal, practices that breach the law occur and the extent to which they occur is not clear.

4.6.6 Potential conflicts of interest in supervisory body membership

Some of the physicians involved in recently high profile, controversial cases in Belgium sit on the Federal Control and Evaluation Commission, raising questions about the sufficiency of procedures to ensure the independence of the key regulatory body.\textsuperscript{131}

4.6.7 Very few officially reported procedural failures

The number of reported procedural failures calls into question the effectiveness of current controls.

In Belgium, as of 2015, the Federal Control and Assessment Commission had only referred a single case of euthanasia to the Crown Prosecution Service out of more than 9,400 cases reported.\textsuperscript{132} In the Netherlands, out of 4,178 cases of euthanasia in 2012, 10 were found to be non-compliant and of 5,512 cases in 2015, 4 were found to be non-compliant.\textsuperscript{133}

A near zero non-compliance rate stretches credulity given the well documented cases above of:

- the existence of widespread practices in the grey area of what is legal, even in countries with legalised euthanasia/assisted suicide, including: publicly reported cases involving deaths that stretch the bounds of the law; and practices involving the use of life-ending drugs without explicit patient request;
- growing numbers of people undergoing euthanasia with mental illness and/or depression;
- regulatory agencies openly conceding their own limitations in monitoring the actual number of instances of euthanasia;
- doctors not reporting cases to the central authorities; and
- other procedural irregularities.


\textsuperscript{133} Dutch Regional Euthanasia Committees, see 2014 and 2015 reports.
4.6.8 Failure of drugs performing as designed

In Oregon, according to official statistics 24 patients are known to have regurgitated some of the legal dose and six regained consciousness after taking the drugs and died later.\(^{134}\)

In Washington State, according to official statistics, there have been at least 8 cases of regurgitation, 2 cases of "other" complications (2012 and 2015) and a seizure (2014).\(^{135}\)

A study of euthanasia and assisted suicide in the Netherlands found that complications arose in 7 per cent cases of assisted suicide and problems with completion (such as a longer-than-expected time to death, failure to induce coma or induction of coma following the awakening of the patient) occurred in 16 per cent of cases. The rate of complications and completion were lower in the case of euthanasia (3 per cent and 6 per cent respectively).\(^{136}\)

The proportion of complications is likely to be higher in assisted suicide schemes (such as that proposed in the Majority Report) than with euthanasia as it will generally be more difficult to ensure appropriate supervision of the administration of the lethal dosage in cases of assisted suicide.

Box 3: Failure of Safeguards

The following safeguards have proved to be difficult to enforce:

- **Reporting**: in Belgium, half or more cases of euthanasia are not reported to the FCEC. This is also a problem in the Netherlands.

- **Patients with depression**: suffering from depression does not mean that a person lacks competence to make important decisions, including whether to opt for euthanasia or assisted suicide. However, it is well documented that serious depression can impact on decision-making, which makes it an area of particular concern given the irreversibility of a decision to hasten one’s death. In addition, the categorisation of some forms of depression as treatment-resistant is problematic as numerous studies have indicated that, in at least some instances, such conditions can be managed.

- **Mentally ill patients**: the number of patients with mental illness being granted access to euthanasia and assisted suicide is increasing rapidly. There is considerable evidence of situations in which properly trained psychiatrists are not being adequately consulted.

- **Coercion**: it is well documented that people who are terminally ill often feel as though they are a burden on their family or carers. This can result in a subtle indirect coercion or, in some instances, a more overt coercion to actively consider a hastening of death.

- **Independent consultations**: in all jurisdictions, there is either a lack of transparency or at least some evidence that independent reviews are not occurring in some cases.

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For the number of instances of regained consciousness see Table 1 of the 2015 report but also, for more detailed breakdown, see Oregon Public Health Division Report for 2012, Table 1, note 13. [http://public.health.oregon.gov/ProviderPartnerResources/EvaluationResearch/DeathwithDignityAct/Documents/year16.pdf](http://public.health.oregon.gov/ProviderPartnerResources/EvaluationResearch/DeathwithDignityAct/Documents/year16.pdf)


4.7 Failure to monitor what occurs outside the law

It is true that many practices in Victoria at present operate outside the full view of the law. But to suggest that all activities undertaken at death are transparently monitored in regimes with legal euthanasia or assisted suicide is clearly not correct.

If anything, it is possible that a normalisation of the hastening of death might create an environment in which even more dubious activity occurs in the grey area that is known to exist but not actively monitored by law enforcement agencies or health regulators.

As noted above, in Belgium, the FCEC has conceded that it has little power to investigate what occurs beyond the reports that it receives from physicians. This is true to varying degrees in all jurisdictions.

Chin et al note that the Oregon Health Division is charged with collecting data under the Dying with Dignity Act and to report cases of noncompliance to the Oregon Board of Medical Examiners. However, they also note that:

Our responsibility to report noncompliance makes it difficult, if not impossible, to detect accurately and comment on underreporting. Furthermore, the reporting requirements can only ensure that the process for obtaining lethal medications complies with the law. We cannot determine whether physician-assisted suicide is being practiced outside the framework of the Death with Dignity Act.\(^\text{137}\)

These examples are corroborated by expert evidence to the Canadian Supreme Court. Professor Etienne Montero was retained by the Attorney-General of Canada to provide impartial, expert opinion to the Supreme Court of Canada in the case of \textit{Carter v the Attorney-General of Canada et al.}\(^\text{138}\) When commenting on the Belgian regime, his analysis led him to conclude that:

the provisions of the Act, as seemingly strict as they are, cannot be strictly enforced and controlled ... \(^\text{139}\)


\(^{139}\) Montero, Professor Etienne, Affidavit to Lee Carter v The Attorney General of Canada et al, Supreme Court of Canada, No. 35591, p41.
Box 4: Legalisation simply shifts where hidden activities occur

The argument that a legalized system creates transparency and ensures a clear understanding of what is happening in real life thus needs to be qualified: it gives us more data, but hidden practices remain and interpretation of the reported cases of compliance and non-compliance call for further refinement and interpretation. Although impossible to obtain reliable evidence, it could very well be that there is simply a data shift, with previously hidden practices now regulated, and an increase of other forms of euthanasia, including involuntary euthanasia and practices that do not respect the legal procedures. The post-factum reporting systems of Belgium and the Netherlands neither prevent this nor provide us more certainty about what is indeed happening. They could just as easily give us a false sense of security and control over the practice. 140

4.8 Noteworthy individual cases

This paper has largely focused on broader, peer-reviewed quantitative studies. This section will highlight some individual cases that relate to concerning trends in the current practice of euthanasia and assisted suicide in some jurisdictions. While focusing on individual cases can be criticised as resorting to anecdote, these cases usefully supplement the studies cited above.

Advocates of euthanasia often cite individual cases so as to stress the human consequences of the restrictions contained in current laws. That is appropriate. It is equally appropriate to stress the human side of the negative consequences of regimes that aim to help a certain group of people but inadvertently result in harm and abuse more broadly.

The following are a small number of the many problematic cases that have made it into the public sphere. It is likely that many more escape public attention:

- **Psychiatric patients:** the euthanasia of a psychiatric inmate in Belgium who was, at the time, incarcerated. Dr Marc Moens, President of the Association of Beige des Syndicats Medicaux (ABSyM) noted that: “Even if the request for euthanasia meets all the statutory conditions, the burning question in this social debate is whether the inmate would have made this decision under the appropriate psychiatric treatment.” 141

- **Transgender:** the euthanasia of Nathan Verhelst (born Nancy) after an unsuccessful sex change operation has raised questions about whether more support could have alleviated the distress experienced after the operation. 142


141 Moens, Dr Marc, president of the ABSyM, press release [translation] Psychiatric inmates have the right to medical care (13 September 2012), and see referred to in Montero, Professor Etienne, Affidavit to Lee Carter v The Attorney General of Canada et al, Supreme Court of Canada, No. 35591, p44.

142 La Libre Belgique (October 1, 2013) and also reference in Lemmens article, pp50-51.
• Anticipation of future pain or suffering: there are a number of instances of people being euthanised as a result of their anticipation of future pain, including Hugo Clause (renowned Flemish author); Christian de Duve (Nobel Prize in Medicine 1974); Emiel Pauwels and others. 143

• Anticipation of blindness: another high profile case was the euthanasia of twins Eddy and Marc Verbessen who were born deaf and were euthanised together at age 45 on the basis of a diagnosis of glaucoma which was gradually making them blind. 144

• Tinnitus: a 47-year old woman in the Netherlands was euthanised for tinnitus that was deemed to be incurable. 145

• Depression: there are many high profile cases where arguably treatable depression was the cause of the euthanasia, including the highly publicised case of Godelieva De Troyer as reported in the New Yorker. 146 In Oregon, the case of Michael Freeland received public attention as he consented to the release of his records. Over a year after receiving his prescription, Mr Freeland was admitted to a psychiatric facility with depression and suicidal intent. His treating psychiatrist wrote a letter to the court the day after his discharge saying he was not competent and needed a guardian. Mr Freeland accidentally called Physicians for Compassionate Care (he was trying to contact the suicide advocacy organisation Compassion in Dying) and was treated for his depression and assisted in reconciling with his estranged daughter. Several weeks later he died naturally and comfortably without taking lethal drugs - some two years after receiving his first lethal dose prescription. 147

• Multiple disorders: the case of Jeanne is illustrative. Jeanne was 88 years old and suffered from multiple conditions that cumulatively caused unbearable pain but that were not individually sufficient to justify euthanasia. For her former attending physician, it was "obvious that she did not have a serious incurable disease as required under the Act". 148 The number of cases of "multiple disorders" increased from 3 in the FCEC's first report to 57 in the fifth report. 149

• Old age: the case of Amelie Van Esbeen gained considerable publicity as it appeared that her old age was the principal reason for her being euthanised. Of note was that her attending physician refused to grant her request.

• Doctor shopping: The case of Kate Cheney in Oregon outlined above in detail is a good example of doctor shopping. The opinions of more than one doctor were essentially ignored. 150 Another instance of doctor shopping in Oregon involved a patient of Dr Charles Bentz, who would not prescribe a lethal dosage as the patient had documented depression for which he needed treatment. The patient's oncologist found another physician required to provide the "second opinion" and did not refer the patient back to his primary physician at all. Dr Bentz obtained

143 Montero op cit pp 45-45 and see also discussion in Lemmens, op cit, pp51-52 in relation to the elderly couple Anne and Francois, who were suffering from significant medical issues but were not at immediate risk of dying. They requested a desire to be euthanised together to avoid the prospect of being alone should one survive the other.

144 Montero op cit pp 45-45 and also discussion in Lemmens, op cit, pp49-50.

145 See this reported in the Guardian: http://www.theguardian.com/lifeandstyle/2015/sep/11/assisted-dying-dutch-end-of-life-netherlands-unbearable-suffering

146 Avid, Rachel, "The Death Treatment", The New Yorker, June 22 2015.
See: http://www.newyorker.com/magazine/2015/06/22/the-death-treatment

147 See information provided by Vermont Alliance for Ethical Health Care, www.vaeh.org.
See also: http://www.pccef.org/articles/art60.htm

148 Montero, Professor Etienne, Affidavit to Lee Carter v The Attorney General of Canada et al, Supreme Court of Canada, No. 35591, p46.

149 Montero, Professor Etienne, Affidavit to Lee Carter v The Attorney General of Canada et al, Supreme Court of Canada, No. 35591, p49.

permission from the patient's family and obtained a copy of the death certificate which stated that death was from malignant melanoma.\textsuperscript{151}

- **Feeling like a burden:** The case of Kate Cheney is an example of the potential impact of pressure due to feeling like a burden on one's family.\textsuperscript{152}

- **Difficulty in diagnosis:** In Oregon, Barbara Houck, aged 71, was diagnosed with Amytropic Lateral Sclerosis. She met with Dr Peter Rasmussen, an oncologist (but not a neurologist) and supporter of physician-assisted dying. Dr Rasmussen declined to give the prescription because he judged that she had more than 6 months to live. She saw him again in March and he gave the prescription. This case raises difficulties about the suitability of an oncologist giving an evaluation and prognosis for an uncommon neurologic disease.\textsuperscript{153} Another case involved a cancer patient, Jeanette Hall. A doctor had given her a terminal diagnosis of six months to a year to live, based upon not being treated for the cancer. She was referred to Dr Kenneth Stevens. On her first visit with Dr Stevens she made it clear that she did not want to be treated. After suggesting that she discuss the idea with her son, she agreed to treatment and, at the time of Dr Stevens' affidavit, had been alive for 13 years. In his view, "the mere presence of legal assisted suicide had steered her to suicide."\textsuperscript{154}

4.9 Conclusion

This chapter has explored various reasons why it is likely that the number of instances of euthanasia and assisted dying continue to grow so strongly, even after a period of almost two decades following legalisation in some jurisdictions.

The most plausible explanations for the growth in overall numbers are:

- An increase in the scope of regimes, whether by legislative amendment or the reinterpretation of existing legislative provisions.
- A normalisation of the decision to opt for euthanasia or assisted suicide.
- An increase in the number of cases of euthanasia of people in a vulnerable position.
- A failure of safeguards to limit the operation of schemes within the boundaries originally intended.

The evidence of Professor Etienne Montero to the Canadian Supreme Court summarises the risks that legalisation would expose Victoria to:

 legislators openness to euthanasia inevitably leads to certain abuses and excesses, to a violation of the letter and the spirit of the law, and to a broadening of the scope of the Act beyond the borders initially and firmly established.\textsuperscript{155}

\textsuperscript{151} Vermont Alliance for Ethical Health Care at www.vaeh.org.
\textsuperscript{153} Vermont Alliance for Ethical Health Care, www.vaeh.org.
\textsuperscript{154} Stevens, Dr Kenneth R., Affidavit in Montanans Against Assisted Suicide & For Living With Dignity vs Board of Medical Examiners, Montana Department of Labour and Industry, sworn at Craig D. Charlton Charlton Law Firm, 16 October 2013.
\textsuperscript{155} Montero, Professor Etienne, Affidavit to Lee Carter v The Attorney General of Canada et al, Supreme Court of Canada, No. 35591, p41.
CHAPTER 5 – ISSUES WITH THE PROPOSED REGIME

The proposed regime for assisted dying contains a number of elements that have proved highly problematic in jurisdictions with legalised euthanasia or assisted suicide. While the scheme set out in the Majority Report claims that it contains a number of safeguards, there is serious doubt as to whether these will be effective in practice.

These include:

- **Scope creep.** Given what has occurred in other jurisdictions, the potential for any entity that is established to expand the scope an assisted suicide regime in Victoria through non-transparent reinterpretations of legislative provisions should not be discounted.

- **Independent review.** The proposed regime would do little to prevent doctor shopping, a practice that is widespread in other jurisdictions. This calls into doubt the effectiveness of the requirement for independent appraisals.

- **Post-factum reporting.** This approach to reviewing cases has proved to be ineffective in a number of jurisdictions, with low rates of reporting and transparency in Belgium and the Netherlands.

- **A requirement that patients have a terminal condition.** As is noted above, this is extremely difficult to enforce in practice, with doctors often disagreeing on whether a condition is terminal. This will be exacerbated by the fact that the requirement will not contain a specific time limit. Even in jurisdictions with a time limit, it has been difficult to enforce this condition.

- **That physicians will be able to administer drugs in certain circumstances.** This is a significant expansion from some assisted suicide regimes. Given that many aspects of regulatory environments have expanded in scope through interpretation, this could become a major loophole.

- **Psychiatric evaluations.** The proposed regime does little to deal with the growing problem of how to diagnose and treat depression and mental illness and, in addition, how to interpret and manage the impact of such conditions on consent.
CHAPTER 6 – ISSUES WITH POLLING RESULTS

It is often claimed that a strong majority in the community support euthanasia or assisted dying. It is true that a number of polls have been undertaken over the past two decades and that somewhere between 65 and 80 per cent of those polled have indicated support for euthanasia.

Given the limitations of some polls to date, it would be worthwhile examining public sentiment in a more detailed, rigorous manner.

6.1 Issues that warrant further examination

There are a number of methodological issues with of the polls that are cited by euthanasia advocates:

- The question is sometimes put in a confusing or ambiguous manner. For example, terms like “dying with dignity” and “assisted dying” could mean different things to different people. Some people consider the use of high doses of pain relief, even if it might have negative consequences, as “assisted dying” – yet this is clearly legal at present and considered part of standard practice by the medical profession.
- At least some of the polls suffer from “framing” issues. It is well established that the way a question is framed can affect the results. In the case of euthanasia and assisted dying, support for legalisation could potentially be affected by references to whether safeguards are seen as appropriate and whether the person being polled would be concerned if was established that safeguards are not working in at least some jurisdictions.

A more rigorous approach would be to undertake layered questioning that disentangles questions such as withdrawal of treatment, the application of pain relief (even if it might hasten death) and euthanasia/assisted suicide.

It would also be worthwhile undertaking polling with groups who have been informed about current best practice palliative care options. It is well established that providing balanced information to people prior to polling can have a material impact on results.

6.2 Specific wording of recent questions

It is worth focusing on some of the specific questions that have been asked in recent polls.

Ambiguous wording

In 2014, the ABC Vote Compass found 76 per cent support for the following proposition:

Terminally ill patients should be able to legally end their own lives with medical assistance

This is a good example of a question that is arguably ambiguous. The phrase “... with medical assistance” could mean the administration of high dosages of pain relief in the eyes of many—a practice that is currently permitted under the law. It is important to be clearer than this proposition.

**Support for tightly defined entitlement**

Where there is support for a clearly defined proposition, it is almost always confined to people who are terminally ill and experiencing unendurable suffering.

In 2015, there was 72 per cent support for assisted suicide in a poll commissioned by Essential Media Communications. The question was:

> Q. When a person has a disease that cannot be cured and is living in severe pain, do you think doctors should or should not be allowed by law to assist the patient to commit suicide if the patient requests it?\(^\text{157}\)

This question includes the phrases “cannot be cured” and “living in severe pain”.

In a 2012 Newspoll, over 80 per cent supported the following euthanasia in the following situation:

> hopelessly ill patient, experiencing unrelievable suffering, with absolutely no chance of recovering

As is discussed in detail above, many people currently opting for assisted suicide in North America and euthanasia and assisted suicide in Europe are not in severe pain at the time of the request. Moreover, in assisted dying regimes of the type supported by the Majority Report, there is little evidence that lethal doses are necessarily only given at such times.

It would be worth testing support for the proposition if it was in a broader context. In particular, it would be worthwhile testing community support for assisted suicide and euthanasia in situations not involving unendurable physical pain, such as anticipation of pain, loss of hope etc, which routinely give rise to assisted suicide in jurisdictions where it is legal.

### 6.3 The need for layered polling

It would be worthwhile testing framing effects. For example, is public support for assisted suicide or euthanasia contingent on the effectiveness of safeguards? Does the level of support change if people are aware of the rapid growth in the number of cases of assisted suicide and euthanasia in all jurisdictions where it is legal and the large body evidence calling into doubt the effectiveness of safeguards?

In polling undertaken by Environics Research Group, a majority of Canadians either strongly support (21%) or somewhat support (36%) legalising euthanasia. This is considerably higher than the proportion who strongly oppose (20%) or somewhat oppose (8%).\(^\text{158}\)

In the same survey:

- almost two thirds of those surveyed expressed concern that elderly Canadians could be pressured into accepting euthanasia (66% concerned vs 33% not concerned).\textsuperscript{159}
- almost eight in ten express concern that a significant number of sick, disabled or elderly could be euthanised without their consent (78% concerned vs 20% not concerned).\textsuperscript{160}

These are not mutually inconsistent positions. But it suggests that, at least to some degree, support for the legalisation of euthanasia and assisted suicide is contingent on the implementation and maintenance of effective safeguards. It is worth exploring how an informed, evidence-based discussion about these concerns could affect support levels.

Abingdon Research undertook a poll of attitudes to euthanasia in Quebec. 74% of respondents either strongly or moderately supported the legalisation of euthanasia.

The people surveyed were then asked to respond to six potential scenarios each dealing with risks associated with the reform. A significant proportion expressed concerns in relation to these risks. After being exposed to these concerns, 47% indicated that they felt the subject required further study, while 35% continued to advocate for legalisation.\textsuperscript{161}

It would be worthwhile exploring the relationship between the risks associated with potential reform and the level of support for legalisation.

\textsuperscript{159} Environics Research Group, \textit{Canadians' Attitudes Towards Euthanasia}, Commissioned Research for Life Canada, October 2010, p11.
\textsuperscript{161} Abingdon Research, Quebec "Medical Aid in Dying Survey", 2013, p25.
RECOMMENDATIONS

RECOMMENDATION 1: That the Victorian government undertake rigorous analysis of trends in euthanasia and assisted suicide in foreign jurisdictions before deciding whether to develop a legal regime for consideration. This would include an understanding of:

- best practice palliative care in Victoria, Australia and internationally.
- the prevalence of failure in safeguards in euthanasia and assisted suicide regimes.
- the extent to which illegal activity continues to occur in jurisdictions with legal euthanasia or assisted suicide.
- the extent to which a rising number of instances of euthanasia or assisted suicide is impacting on the overall health care system.

RECOMMENDATION 2: That rigorous, layered polling be undertaken in Victoria and that this polling:

- clearly set out the scope of the reform being tested.
- adopt a range of different "frames" for the question, including frames that indicate the risks associated with containing the scope of euthanasia or assisted suicide in practice.
- explore the relationship between a person’s awareness of and attitude towards the various risks associated with legalisation and their level of support for legalisation.
CONCLUSION

The legalisation of either assisted suicide or euthanasia would represent a fundamental shift in the regulation of medical practice in Victoria. Overseas experience over almost two decades suggests that, while such a change may benefit a small minority of people toward the end of life, many more may be worse off as a result.

This Minority Report has argued that:

- It is possible to respect individual autonomy while not empowering health professionals to actively participate in acts of assisted suicide or euthanasia.
- Even if it can be argued that euthanasia or assisted suicide are justifiable in some instances, the negative consequences arising from legalisation far outweigh the benefits arising in that minority of cases.

In only a very small minority of end-of-life situations are the symptoms of pain unmanageable

Experts in palliative care, oncology and related fields provided the Committee with powerful evidence that almost all symptoms arising from physical pain at the end of life can now be managed. The palliative care and oncology experts who gave evidence also stated that, over long careers, the number of people expressing a desire to have their life shortened was very small. Moreover, pain relief techniques and palliative care are constantly improving.

Even where there is an expressed desire to die, it is critically important to understand the nuances of such requests. Where the person making the request is experiencing depression or a mental illness, which is relatively common, there are usually treatment options worth exploring. Holistic palliative care and other forms of assistance can often provide effective relief, even if not complete, and can often lead to a reversal in the expressed desire.

Consideration of assisted suicide and euthanasia requires a consideration of more than how best to give effect to individual autonomy

Proper consideration of assisted suicide and euthanasia requires an evaluation of the likely broader social impacts of the reform in addition to issues relating to individual autonomy. Individual autonomy is of great importance, but it is not the only consideration.

While there is arguably a gain from giving effect to some patients’ wishes to hasten their death, this must be weighed against the potential for societal harm through a range of potential negative consequences, including: vulnerable people being pressured into euthanasia or assisted suicide; people having their death hastened without having given proper consent; and a gradual broadening of practices without transparent public consideration.
In practice, euthanasia and assisted suicide are a disproportionate response that cause far more social harm than good.

The evidence is clear that the number of instances of euthanasia and assisted suicide is growing rapidly in all major jurisdictions where it is legal. This has been occurring for almost two decades in some jurisdictions, with no sign of abatement.

The usage of euthanasia and assisted suicide in practice is far out of proportion to the situations that were originally used to justify the practice in these jurisdictions: namely, that small minority of cases where the symptoms of pain are unmanageable. Moreover, the rapid growth in documented cases of euthanasia and assisted suicide probably materially understates the actual prevalence of the practice given low rates of reporting.

The effectiveness of safeguards in jurisdictions with legalised assisted suicide and euthanasia has been called into question by many academic studies and high profile media investigations. This, coupled with the growing prevalence of the practice, is a matter of great concern.

While legalisation was supposed to bring what was occurring in the shadows into the light, legalisation has simply pushed the boundary of what is legal out further and may have increased the amount of activity that occurs beyond the sight of regulators. Evidence suggests that it is doubtful that safeguards are working as intended, particularly for vulnerable people.

It is very unfortunate that some symptoms of pain cannot be totally managed— but the legalisation of euthanasia or assisted suicide is not the appropriate response.

Daniel Mulino

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If you want to change society, the first thing you have to do, all social engineering is preceded by verbal engineering. That's one tactic. So do not let any of these euphemisms for assisted suicide be used over and over again.

The second leg, if you can't fool people with just language alone, you have to create fear.

Now I'm going to share facts with you, and when I share facts that sometimes are kind of sad, people say that's causing fear. Well if it's true, you should know about those facts. But if it's not true, then you need to basically create fear that you're going to be suffering, in great pain, and the only way to relieve your suffering, the only solution to suffering, is to end the life of the sufferer.

That's the second tactic. And when you allow reasoned dialogue like this in legislatures here or elsewhere, generally the assisted suicide movement loses. Because as much as we like to bash politicians, they have hearings, as you all know who are here in Parliament, sometimes for hours, days, months, and there is a time to go past drive-by debate, and generally when you look at the facts, as Margaret said, worldwide, it's not pretty.

So if you can't win with reasoned dialogue, then you go ahead and shift the issue to faith, and church, and you attack the church or the faith of the individual who's espousing the reasonable arguments that make this not a good fit for society.

So those are the three legs of the stool, euphemisms, fear, and don't talk about the subject but talk about the fact that I might have faith or I might not have faith.

One of the more powerful spokespersons for the opposition to assisted suicide in society is a secular humanist lawyer in Seattle, Margaret Dore, who has a website that I'd encourage you to look at if you want facts about this. It's called Choice is an Illusion. She resents the fact anytime brings up issues about overall life or death or choice. She wants to talk about the facts, as many
times, lawyers and barristers and judges do, just the facts. And she'll chastise me if I drift apart from that topic.

So I chose the title of this little presentation, I made it a little terser than the ones I've given in the last two days, because I believe your time's valuable, I want to give you plenty of time to talk. If I don't make something clear, please feel free to interrupt me. I didn't title this "The 10 Myths", I titled it "10 Myths" because there are more myths about assisted suicide. But I want to go through those things, and my goal with this is to show that when you exercise this so-called right, in Oregon or Washington, you are displaying dispassion rather than compassion, and I'll show you some specific examples that illustrate that.

Now don't get me wrong, I'm not attacking people individually, but I'm telling you when you're apathetic about what people choose, it isn't being willing to suffer with, that's the roots of the word, it's being dispassionate. It's colluding with that feeling.

I want to mention that it's dangerous because there have been cases where people are going after ulterior motives. Not everyone is a Boy Scout or Girl Scout in the world, and certainly not in medicine. In the U.S., there are over a million doctors. They all don't have the same ethic.

It has numerous problems, I'm going to only in the brief time I have, highlight some of them. And there are alternatives. It's Physicians for Compassion and Care. We chose the name deliberately. We're not an opposition group. We're trying to enhance end-of-life care, but never compromising our integrity to embrace situational killing at the end of life. We have the technology and the capacity, we have the humanity to do that.

In Australia, I've read in your paper where you're cutting back on home deliveries, not home deliveries, it's home care out of cost saving. In just yesterday or the day before's paper, Margaret talked about how things are getting cut back because in Australia, like in the United States, costs are going up in health care faster than the rate of inflation.

**So Myth Number One, you'll hear this over and over again of it's because of pain.**

Actually, pain is seldom the reason, and I'm going to show the data that supports with I'm contending. If anything, it's the fear of pain.

Now I agree, there is fear of pain. I'm not just talking theoretically either. You know, I've lost five of my closest loved ones in the last 15 years, beginning with my father in 2002, 2003, my mother, three and a half months later. My mother and father-in-law had a brain tumour, and she followed six or seven months after my father-in-law died, and 37 months ago, my wife of 40 years, I lost.

All of them died with dignity. None of them took an overdose. So I'm not talking theoretically. I'm talking in real terms of applying the facts and the information I'm sharing with you. It is true that virtually all pain can be controlled.

My father, after an emergency colectomy, got a propofol drip. That's where the ICU nurse administers intravenous medication to keep you sedated so you don't fight the respirator, you don't pull out tubes inadvertently. And by the time they lightened the drip to bring him back to consciousness, he had no pain whatsoever. I had to tell him what went on because he was doing so well.

The sedation was used as a technique to control his pain, make him comfortable, with a goal of having him be more robust as he recovered from surgery. So we have that technology, it was used on my own family.

We can use radiation, my wife got this for some of her cancer, trying to hopefully help the pain in her spine. When that wasn't totally effective, she had what's called palliative surgery. Palliative
surgery is where that metastatic fracture of her spine, that's where cancer goes into the bone, weakens the bone, the spine collapses, is threatening to make her a paraplegic for the last few months of her life. And because we had private insurance, the palliative surgery made her able to walk without pain. The surgery was nothing compared to the pain that she had before it.

We attended our son's graduation two weeks before. He graduated from the University of Dallas, which was a life-changing experience for my son, who still talks about how special that was. That would not be possible if we were covered by the Oregon Health Plan, which would restrict useless surgery, because she's going to die anyway.

Now I'm not making this up, I'll show an actual case of this. And again, people will say these are anecdotes. These are the things that have leaked out from the shroud of secrecy in the state of Oregon, because the state doesn't report this kind of thing.

There's a shroud of secrecy. What I mean by that, you have no capacity as a citizen or a reporter to go into the details of what the government's recording. The information given to them is all second and third-hand, because the doctor's not present 86% of the time. So who really knows what's going on? And if you're not bothered by that, the records are actively destroyed within a year after they've collated the data.

There's no ability or authority or finance for the Oregon Health Division to look into the records. And yet you'll hear over and over again, if you haven't already heard it, everything's going swimmingly in Oregon.

46% of the time, the space for complications on that form the doctors are supposed to fill in doesn't say none, it's blank. It has to be blank, because if they said none they would be lying. They don't know what happened, they weren't there.

There's an inverse relationship between pain and the desire for assisted suicide. It's the fear of pain.

Am I making this up? No, this is published in the Lancet, 1996, it's been known for a long time. As people actually experience pain, and we are able to work with them in controlling it, their desire for suicide actually drops, as is displayed in this graph outlining what the article showed.

The desire for suicide is on the vertical axis. And experiencing pain up to 10 on a scale of 10, is on the horizontal axis. While the data is flawed, as I said, what data we have supports what I just said. It's not pain, it's losing autonomy or so-called dignity.

Now I believe in death with dignity. I believe in aid in dying. I believe in choice in dying. I believe in options at the end of life. If that's what we're talking about, we have no debate. I don't believe in empowering doctors to end the lives of patients with massive overdoses or with injections.

Can we be clear about the king's English? Can we break through the euphemisms, and be honest about what we're talking about? Because currently, people have the right to kill themselves as was shown in the Queensland just yesterday, or the day before. I was talking about it with the press yesterday. I think it's a tragedy.

I think they were enabled by Dr Nitschke, who sells canisters of nitrogen because apparently the helium people don't want to be in the business of killing. And I think it's a tragedy, and I think it's good that the reporter put at the end of it, if you're having suicidal thoughts or you're depressed, here, call this line. I think that's the way society, I think that's the way Oregon should, and Australia already is, helping people who might have those thoughts. It's trying to avoid suicide contagion.

Unfortunately, in Oregon, we've dropped the money for suicide prevention, now the only suicide prevention we have in Oregon is for adolescents. Apparently, once you get past 21 you're not as
valuable as are young people is what the message we're saying. But what are we telling young people when we tell them the solution to suffering is to end life, to end your own life.

In the U.S., we just had somebody indicted and actually convicted of aiding somebody's suicide, when online relationship, she actually encouraged him to go through it and gas himself with carbon monoxide. And appropriately, she's being charged with manslaughter. But this whole notion of thinking it's okay with your radical autonomy to do this is dangerous, and that's what I said at the outset. I'm trying to show you this has got consequences besides the private issue of the individual.

By the way, look at the other reasons. Decreases in the activities that make life enjoyable. Loss of dignity, again there's that term.

This is an insult to people who are disabled with incontinence, with paraplegia. There is a group called Not Dead Yet, who recognises the threat to their well-being with this paradigm. Now think of this, men obviously have difficulties with obstruction of their bladder sometimes because we got too much anatomy between us and the outside world. As we age, that becomes more of a problem. Almost all my mine over 65 or 70 have changes in the way they urinate. Women, on the other hand, have a different problem. They don't have enough anatomy between them and the outside, and they actually step off a curb, you sneeze, you laugh, whatever, and so is this making you less dignified? I'd like to see a show of hands. No. And yet that was raised by Judge Goodwin in a 110-page opinion in the Ninth Circuit Court that that might be a reason, beyond being terminal, to have assisted suicide.

The Ninth Circuit Court. Fortunately, it's one of the most overturned courts, federal courts in the United States. But that kind of reasoning is dangerous is my point. You see what I'm saying?

Myth Two, there are no problems in Oregon.

I've already mentioned the reporting system's flawed, the doctor's not present 86% of the time, it's second and third-hand information. There is never any investigation by the Oregon Health Division. This is a quote from Katrina Hedberg herself. "We don't have the authority nor the finances to investigate."

So if you are going to go there, and I'm not recommending it, ask what they're going to do and who's going to be doing the investigation, who's going to pay for it, who's going to triangulate, do the morbidity and mortality, to be sure that things were aboveboard? Who's going to be the witness at these things to be sure it's happening correctly? Because they say it's so safe.

Is any procedure without problems? The UK Select Committee, which was much like your committee that's gone around and looked at these issues, had the wisdom, because they had at least two doctors, a palliative care doctor and a surgeon, and the surgeon on that committee said, Any time I hear someone say there's no complications with a procedure, I know they're either not aware of it, they're ignorant, or they're lying. I'm paraphrasing him.

Do you know people who can take an aspirin always with no problem? And that's what's being said. For 20 years, the promoters of assisted suicide have repeated this untruth over and over and over again. So do not allow the ministers who are in favour of this to keep reciting this untruth.

Often the reports that are given, of course, are not only by doctors who engage in it, but people who actually promote it. I know because I work with some of them. I know the doctor who actually gave Brittany Maynard her overdose. He works in my department. He works in palliative care. I know at least three or four doctors in my own department who participate in this, and I know some of them have been featured in television commercials, some of them have written articles about it in peer-reviewed literature. So you're asking the very people who are in favour of this to tell you
that there are problems with it. And that's a little bit like asking the Tobacco Institute to say what's wrong with tobacco smoking.

In fact, as I said, the blank spaces on the complications are on 543 out of 1,127 reports turned in by doctors who are participating in this. That's almost half, 48% of them are blank. Again, that's actually honest because they weren't there, how could they fill it out? We really don't know what's happening, yes, yes.

[Man] In the Oregon system, do you need two doctors? Do you have to get a companion to support? The question is, in the Oregon system there are two doctors. Do you have to get a companion to, yes. The way, that so-called safeguard is, you have to agree that it's your decision that this is a good idea for that patient, fitting the criteria, and you have to get a colleague to say yes as well.

It can be any doctor. It can be a pathologist, it can be a paediatrician. It can be a psychiatrist, it can be a palliative care specialist. It could be someone who knows nothing about anything about end-of-life care. And yet if they have an MD or a DO, or in the UK it would be an MB, whatever, you're okay to do this.

And there's a network now, and it's fueled by so-called Compassion and Choices, the group that's promoting this worldwide. They have a network of doctors, and they're all too eager to help you connect with a doctor, who also knows other doctors who will be a second signatory. When somebody doesn't use these known doctors, and a doctor refuses, they simply keep shopping for another doctor.

**Myth Three, you have only six months to live.**

My wife was told we have three to nine months to live, two or three years before actually died. She thankfully lived for almost four times that long. Every day was precious. Our lives in our last five years, while I wouldn't wish cancer on anyone, were actually among the best of our 40 years together. We had great suffering at times, but we also had great joy at times, because we knew every day was special, that we might not have another time to do whatever we were doing, and it changed everything.

I think I had one argument with my wife in the last five years of our marriage. I think that was a very different rate of arguments than we had in the first 35. It was over in about 20 minutes, because we realised how out of proportion our being upset with each other was relative to all the good that we had, and that we had fragile time left. So again, I wouldn't wish cancer or suffering on anybody, but it brought us closer together.

There are no crystal ball reading courses in medical school. The reality is doctors become pretty good, and I say pretty good, when it gets down to the last couple days of life. But we're lousy when we're talking three months. You might as well ask weathermen what's the temperature going to be in three months from now. And you know that'd be pretty silly.

Well doctors don't have any crystal ball reading courses. We're not genies or gipsies, and we're wrong a lot of the time. We're wrong even about the diagnosis sometimes. And I've been wrong about patients, and I regret that I was wrong. I gave up on people sometimes who lived, and every time I see this person 10, 20 years later, I'm embarrassed that I gave up on him. But that's true.

Six months is arbitrary. If you believe this, that six months is okay, it's going to control it, why not seven months? What right do you have to keep somebody with seven months? Or seven years? Because if you believe the solution to suffering is to end the life of the sufferer, then you have no moral, ethical basis on which to deny someone who's suffering for seven years with their rheumatoid arthritis or their lung disease. Or their cancer like my wife had, from having it done,
and indeed that is why in the Netherlands and Belgium, they have opened it up, so literally if you feel that you've lived a complete life now, this is the bill being debated, very different than Australia's, but it's exactly where you go, because it was only legalised in the Netherlands in 2002.

They've been doing it illegally since 1971 to 1973, when the Postma case, was extand. He's a GP in the Netherlands, his wife was a GP, and they injected her mother because of her, quote, suffering. The wrist of his wife was slapped. She was given a year in gaol, which was suspended, and she had probation for a year.

And indeed the country then went from that hard case to doing it essentially suicide on demand, where parents in the Netherlands can end the lives of their kids, up to the age of 12, according to the Groningen protocol, where you have people whose lives are ended without permission, there's a category. Over a thousand people a year in the Netherlands have end of life without explicit request.

Now what is ending your life without requesting it? I'll let you describe it. I'm not trying to be pejorative. And when you ask doctors, why did they do that, on what basis did you take it upon yourself to end someone's life? Well, I knew they were going to die anyhow. And I needed the bed. This is published in the Journal of the American Medical Association by Herbert Hendin. I'm sharing with you facts.

Another doctor was taking care of a religious, a nun. "I knew it was against her religion, she was Catholic. So I didn't tell her about it." So much for openness and transparency. And it's over a thousand times every year in two different reports by the government of the Netherlands.

I'm sharing facts, not hyperbole. I'm sharing appropriate knowledge, so you as citizens of Australia, can make informed choices about this coming paradigm.

In Oregon, about a year and a half, two years ago, two of my colleagues who now are legislators, Mitch Greenlick and Elizabeth Steiner, and one other legislator, proposed that we expand it from six months to 12 months in Oregon, predicted life. And both sides were opposed to it. Now you might suspect that someone like me would be opposed to expanding what I believe is a misguided paradigm.

Why would the proponents of assisted suicide be against expanding, since some of them really want suicide on demand? Like Nitschke, like Kevorkian, like Governor Lamb of Colorado. They didn't want to expose, when they're trying to pass it in 25 other states and the District of Columbia, the truth that there is no way to control this. All it takes is a legislative fiat that says, you know, six months is too restrictive.

In Canada, they're already going from physician-assisted suicide to euthanasia. And Quebec's asking all of us to carry not only one euthanasia kit, but a second one for backup because it may not work. At least they acknowledge that these things don't always work. But sadly, that's where we're going.

The ink's hardly dry no the Canadian judges' 9-0 decision that somehow, physician-assisted suicide and euthanasia is contained under the right to life in the Canadian Constitution. You can't make this logic up, but that's what they're saying. Essentially, anyone is eligible, any time for any reason in the Netherlands and Belgium.

The EU is against the death penalty, and I'm not arguing death penalty here, I'm talking about end of life.

So there's a prisoner with a life sentence for rape and murder just in January about two years ago, who thought his life was not worth living because he had no hope of parole and he's in for a life
sentence. So he requested euthanasia, and within a couple months, he's dead. So the same
government that abhors the idea of ending the lives of convicted felons, murderers, rapists,
whatever, allowed him to be killed with euthanasia. There are no abuses.

In Oregon, we've already had nurse-assisted suicide, this was only found out because of astute
reporters in the paper, the Portland Tribune. You can look it up online. They were actually fighting
over this individual, who had had a sex change, was same-sex attraction, and the editors were
wondering about whether they should report these things.

I'm not here to talk about these issues. I'm telling you that there are ulterior motives among some
people in this paradigm. They were not given orders by the doctor, by the way. Opioids are a
controlled substance in every state in the United States. One nurse said we were just doing pain
relief with him. The other nurse said they were doing assisted suicide. All I can say is, one of the
nurses is not telling the truth. This went unreported for almost two years before it leaked out, not
by the government, but it leaked out by an astute reporter.

I mentioned there are no witnesses, so once you get the overdose, it's the perfect murder. Because
you've already been labelled as terminal, you've been labelled as having the right to assisted
suicide, and all the person who's a survivor has to say is that it was an assisted suicide, and you
really don't know what happened because no witnesses, nobody's there. Nobody asks,

I've never asked a patient in my 40-year career, what life insurance policy do you have, and who's
the beneficiary? One out of 10 people in the United States, I don't know if it's true in Australia, one
out of 10 have a problem with elder abuse. The ultimate elder abuse is taking the life of the elder.

There's suicide tourism. This is not me making it up. If all of you have heard about Brittany
Maynard, who had a brain tumour. It was inoperable, it was lethal, there was no cure for it. She
confirmed that at Stanford University, then she decides to become an Oregon citizen and comes
to my colleague, Eric Walsh. I can name names because he said it on 60 Minutes when we were
both on that programme about a year and a half ago, and he was the one who facilitated Brittany
Maynard to have a hastened death with massive overdose of pills. There's doctor shopping for the
right answer. Am I making this up, no. This also leaked out.

The cases I'm sharing with you are not being exposed by the Oregon Health Division. It's not being
controlled because there is no control, but again an astute reporter interviews two people only
because they were frustrated that the system held them back.

On your right is Erica, with the Race for the Cure shirt. And she's next to her mother, Kate Chaney,
who was a survivor of one of the Holocaust prisoner of war camps, or not even prisoner of war,
Holocaust death camps.

Let me correct myself. She was somewhat demented, her mother, and she went to one of my
colleagues who is clearly claiming to be neutral about this topic, and it wasn't clear to the
psychiatrist, Linda Ganzini. I can mention names because all this is published now, who was really
asking for the suicide because the daughter, Erica, seemed to be coercive.

So she goes to a different doctor, and that doctor doesn't use the word coercive, but he also will
not sign the process, begin the process for assisted suicide.

She goes to a third doctor then, who works for a health care capitation system, where you're giving
coverage for everybody for whatever, and yet that doctor thought she was a good candidate for
assisted suicide.

So the capitated system will actually save money by ending the life of Kate Chaney. And indeed
she was dead within a matter of weeks from that opinion. And getting a second opinion is no trouble
because they actually sent a letter around to the Kaiser Group about who would participate in this, and they sent it to 800 doctors in the Portland, Oregon, area to find out who would be a co-signer.

Myth Number Five, death is dignified.

If anyone's worked in an emergency room and seen overdoses, it is not the prettiest way to die. In the Dutch experience, they don't generally do this, they don't recommend it because it doesn't work a lot of the time. In fact, they say somewhere between 17 and 23% of the time they have to be standing by as doctors, and they generally do, with an injection, to be sure they die.

It does cause nausea and vomiting because the pills are extremely bitter. The dying can be prolonged with agonal breathing, with noisy respirations as you're not clearing your secretions from your windpipe.

And again in the newspaper, David Pruitt awoke 67 hours after taking the cocktail. And I'm quoting, his first words were, "What the hell happened? I thought I was supposed to be dead." His wife, who had gone along with him and said, 'whatever you want', trying to, quote, "respect his wishes", was elated that he was still alive. He had an epiphany, feeling like, well, God must want me to live, and he actually had a good next 11 days before he died naturally with dignity. So the co-opting of the word dignity is somewhat, no, it's totally misleading. And the reason why I say that is because my own wife died with dignity. At home, surrounded by family.

And if you say that you have to take an overdose to be dignified, what does it say about the rest of us who don't take an overdose? Implicitly, you're saying something special about killing yourself with an overdose.

Suicide parties are happening. They reflect again, I said at the outset, so-called compassion by taking overdose is really apathy, and I'll share with you that in a minute here.

This is a picture that was on the Oregonian website, OregonLive.com, and if I were to just show you this picture, you would not know if anybody's really sick here. These two people are having a very good time.

The one on your left, though, is Lovelle Svart. And she died in 2007, after having for months been on a video log, posted by the Oregonian, because she used to work for the Oregonian five years earlier, showing the deadly potion she was ultimately going to drink, and she schedules this suicide party in her home. And they're doing the polka, and they're obviously having a good time, and if I didn't give you this background, you'd say these are two people who know each other well and care about each other.
And the next scene, within 30 minutes, is in her bedroom, where you see her taking the deadly overdose of barbiturates, and next to her is not her husband, not her doctor, not a loved one. It's George Amey, who's a lawyer, who's the executive director for the so-called Compassion and Choices.

He's already done, at this point, at least 30 of these, where he's become expert, and he's now being sure that she wants to go through with it, and he says, and I'll paraphrase the conversation. He says, "Now, you're sure you want to go through with this?" She replies, "I'd really rather go on dancing". To which he replies, "Well, you can." She says, "No, I've already taken the antiemetic to keep from vomiting, and we were all planning this", so she has this pressure to go through with it. And she does, and she starts drinking the overdose.

And he coaches her, telling her not to drink it too fast because if you drink it too fast, you're more likely to vomit it and not be successful. If you drink it too slowly, you might fall asleep before you take a deadly overdose.

And then he becomes very dispassionate and clinical in describing her reduced respiration to the rest of the people in attendance, like you're watching a dog lab and sharing with medical students what's happening to the dog's breathing and pulse.

Now what if George Amey, instead of doing what he did, which was a vending machine, well, you can go dancing polka or not, what if he said, "Lovelle, I've come to know you in the last few weeks, and you're really a special person. If you want to go on dancing, that's a good idea. Don't worry, we can give you this sometime in the future, but you've got a lot of life left in you, don't do this. I care about you. In fact, I've come to love you."

Now which of the two responses, well, you can, or the one I just described as a possibility, is the compassionate response to one another as humans? Which is encouraging that you're worthwhile even when you don't feel like it, and maybe you're wrong-headed in feeling your life has no value, or you're fearful about being able to get it?

Which is the compassionate response, I ask you. And what do you want for Australia in the future? What kind of response do you want from your doctors, because I'm a doctor, and what kind of response do you want from me?
Do you want me to be a vending machine? And if Australia wants to embrace this, I want you to know that I would not be in favour it, but I want it not to be in the House of Medicine.

Perhaps you should have veterinarians, who have training and skill in giving overdoses to living things, living animals. I'm not recommending that, I want to be sure, don't misquote me. But at least then you'd keep the House of Medicine from an inherent conflict of interest. And you'd have trust in your doctors.

It's the most profound secular argument, and I heard from the sitting dean at the time, of Oregon Health and Science University, because he's very clear and to the point about the inherent conflict of interest for medicine.

**Myth Six, it improves end-of-life care, and look what's it done in Oregon, it's gotten much better.**

Well the perception of the adequacy of pain control was actually studied by one of my colleagues and published in a peer-reviewed journal. He's a palliative care specialist. His name is Erik Fromme, F-R-O-M-M-E. He actually took care of my wife, and I selected him specifically because I knew he did not promote assisted suicide. I want doctors like that for me and my family. What kind of doctor do you want?

He published a study that said family's perception of pain at the time that it was studied, before and after assisted suicide, perception of pain in their loved one, their dying one, was worse after passage of assisted suicide.

Now I'm intellectually honest enough, I'm not blaming assisted suicide. Because you cannot do that. That's not cause and effect data. It's epidemiological data, but it's not reassuring, and it does not suggest that pain relief has improved in the state of Oregon. Do you follow me?

I was misquoted by that also by one of the promoters of death here in Australia years ago, because they're so bent on the agenda that they will not listen to reasoned dialogue. And you did not hear me say that assisted suicide caused this, but you did hear me say, it is not reassuring in Oregon that this is improving pain relief.

And in fact the opioid use per capita in Oregon was high well before the passage of assisted suicide. It's still high, but it's even higher in states that have specifically made it illegal to have assisted suicide.

There's no correlation between passing appropriate laws that circumscribe the power of physicians to support an enhanced life until you die naturally; to extend life, and circumscribe it so you never cross that boundary to actively end lives.

By the way, opiate prescribing per capita is actually not a great marker for good health care at the end of life because there's such abuse of it that almost every state now is trying to limit opioid use because we've been so seduced by a doctor back in the '70s and '80s who thought that we could give opioids successfully to treat pain in non-terminal patients. And now we've found that we are killing more people with too many opioids in people who aren't terminal that we are actually having more people's lives end than on highway accidents in the United States.

But it is a marker and a surrogate when you say, oh, people are going to be fearful about giving appropriate doses of opioids. That's false, that's not true. Even in states that have made it illegal to do assisted suicide.
Myth Number Seven, it expands patient choice.

As I said, patients have the right to end your life now, it's not illegal in any country that I know of. There are 43,000-plus that happen in the United States. Margaret alluded to every three hours. It may even be higher than that with the math I would do, but the point is that we have a problem. Houston, we have a problem.

Suicide, non-assisted suicide, is all too common. So you say what difference does it make? We're only talking, Dr. Toffier, about the limited few numbers that are happening in Oregon, Washington, and it'll only be a few numbers here in Australia. This is the data.

One of the ministers said they wanted to not talk to me because they want to do evidence-based discussions. This is the data.


The suicide rate in the bottom is the United States. It's gone up slightly. The blue and the red are both Oregon and Washington. They were above the national average before a passage of assisted suicide, and the slope of those lines is not reassuring.

Again, you're not hearing me overstate the case, saying that assisted suicide caused this, because that would be beyond the evidence. But it's not reassuring. And you know there's suicide contagion.

That's why the reporter told me, "oh, we always have to put that thing at the end of the story about the three women in Queensland who took their lives. We always have to do that." Why do they have to do it? Because they're fearful of suicide contagion, of spreading Dr. Nitschke's false approach to suffering. They're fearful of that, appropriately fearful of that.

And we all should be. And you should increase your spending on suicide prevention, not on assisted suicide. And it is spending by the way.
California allocated $2.4 million for MediCal, that's the welfare system, so they could be sure they
could cover the cost of the drugs to kill yourself. So where did that $2.4 million come out of? Was
it out of prenatal care? Well-child care? Well, their math in their budgetary session to legalise
assisted suicide in the state of California, obviously is that we'll save money by spending $2.4
million to end the life of the sufferer rather than paying for the expensive drugs that might come
forth. So in truth, it's not empowering patient choice.

It's not rocket science to kill yourself. As I shared with one of your ministers, I was in aerospace
engineering before I went into medicine. And it's not rocket science, I didn't take any courses in
this. I don't recommend it.

Patients have the right to refuse treatment. At one time, medicine might have been guilty, and I
came into medicine in the '70s, of doing things that were not reasonable relative to the overall
situation.

Today, that's not the problem. The problem is accessing care. Patients who desire coverage have
been denied coverage, and I'll share at least one of those stories here today.

Barbara Wagner is a 64-year-old woman. At the time, she had a recurrence of her cancer, she
went to her oncologist, her cancer doctor, who wanted to give her the drug Tarceva. Tarceva's a
drug that now has been around for 15 or 20 years, it doesn't cure the cancer, just like the six drugs
my wife took before she died wouldn't have cured her. We were hoping that it would extend her
life.

Tarceva has statistically a 45% increased chance of keeping the person alive and being alive in
one year, if you take it, versus not taking it. Kind of like a woman with breast cancer, if you take
Tamoxifen, you're more likely to be alive than if you don't after you have your breast surgery. Same
kind of deal, doesn't guarantee a cure.

She was given a letter, which went to at least one other person, we don't know how many because
the Oregon Health Division will not tell us. It's under a shroud of secrecy. But at least two people
we know got this letter. It said, we cannot cover your chemotherapy that you want, your oncologist,
your doctor wants, but we will cover 100% for your assisted suicide, under the euphemism of pain
relief and relief of suffering. In the same letter, it says this.

So let's hear from Barbara and what she has to say.

[Reporter] Her doctor offered hope in this bottle, the new chemotherapy drug, Tarceva.

[Barbara] Like my doctor said, maybe this can put the lid on it and stop it.

[Reporter] That hope shattered with this letter from the Oregon Health Plan, telling her, We were
unable to approve the cancer treatment. It will pay for comfort care, including physician aid-in-
dying, better known as assisted suicide.

[Barbara] I told them, you know, I said, Who do you guys think you are? To say that you'll pay for
my dying, but you won't pay to help me possibly live longer.

[Reporter] We took her concerns to Dr. Som Saha, chairman of the commission that sets policy for
the Oregon Health Plan. She says, to say to someone, we'll pay for you to die but not pay for you
to live, it's cruel.

[Representative] I don't think anyone's saying that. I don't think anyone's saying that. That's, I think,
maybe an unfortunate interpretation of the letter.

[Reporter] Assisted suicide critic Dr. William Toffler calls the message disturbing.
[William] People deserve relief of their suffering, not giving them an overdose.

[Reporter] And he says the state has a financial incentive to offer death instead of life. Chemotherapy drugs like Tarceva cost $4,000 a month. Drugs for assisted suicide cost less than 100. Is it cheaper to pay for somebody to die than to help them live?

[William] That is a not a question that we think about. We don't think about, we don't think about investing our health care dollars in that way.

[Reporter] Yet when thinking about patients like Barbara Wagner, Dr. Saha admits they must consider the state's limited dollars. If we invest thousands and thousands of dollars in one person's days to weeks, yet we are taking away those dollars from someone.

So look at the body language and the uncomfortableness of that man in trying to defend the indefensible. Often, people say, gee, these numbers, you got to look at it realistically. $4,000, well Tarceva hasn't changed that much. Suicide pills have gone up greatly in cost. And in fact, the pills that used to cost $100 are now over 3,000. Why? Because the drug companies don't like manufacturing these drugs for death penalty patients, and they don't like to be associated with this. And so there's only one making the product, and they're charging a lot for it.

So the suicide promoters are actually actively looking for other cocktails in which people can kill themselves. Just as Nitschke is no longer using helium, he's using nitrogen. They're working overtime to try to find massive overdoses, so they're suggesting that you take a thousand times the normal dose of morphine and two other pills to be sure you do yourself in. This is sad research, I think, and that's what they're doing.

You say, well, Dr. Toffler, that's just an anecdote, that's in Oregon. Well, in California, it only passed a year ago today. The ink is not hardly dry, and look what's happening in California.

[Woman] I am a wife and a mother of four, and I've been diagnosed with a terminal illness. We've been trying to get me on a different chemotherapy drug for the infusions. And when the law was passed, a week later, I received a letter in the mail saying that they were going to deny coverage. They didn't want to pay the money for it. On my letter, it didn't say anything about physician-assisted suicide, as far as you know, why they were denying me. So I called the insurance company, and I was asking them why I didn't get approved for this drug. So I said, well what about the drugs that they're using for the new law, you know, this physician-assisted suicide, there are these pills that I could get. Would you cover that for me? She says, yes, we do provide that to our patients. You would only have to pay a dollar and 20 cents.

You'd only have to pay a dollar 20 cents is what she said. 80% of the insurance companies in California now cover assisted suicide, paying for the drugs. Yes.

[Man in audience] “Excuse me, Dr. Toffler, are you saying that now, though, the price of the drugs has gone up so much that really the comparison is the same between treatment and the cost of the suicide drugs?”

So the question is, the cost of the suicide drugs have gone up so much that the comparison is the same between the two. I'm hearing the question correctly?

[Man] “Yes, so that governments perhaps won't be influenced in the way that you're expressing.”

So the implications are the governments wouldn't be influenced because it's a wash, it's six of one, half a dozen of the others? Wrong, because, when I say $3,000 a month, $4,000 a month, one of my wife's drugs was $8,000 a month, it's per month. The overdose, even if it was $3,000, is a one-time expense.
[Man] “Thank you.”

Caring is more costly than killing. I believe in caring, not killing. I believe in caring, not empowering doctors to kill people. And if society wants to go down that path, I would encourage Australia to licence veterinarians to do the deed, because they have the most expertise.

I often in the past have asked that question, and I get a blank stare. Well, who would do this? Well, they have some expertise. Now I think, if I were a veterinarian, I wouldn't do that, because I think humans are different. And I think we're a different species and deserve to be treated.

Now, some people might debate that as well, but I do think there's a solution if society believes this should be, I think you should avoid compromising the integrity of your doctors.

**Myth Number Eight, patients are screened for depression and mental illness.**

Actually, that's not true. Is it just my perception? No, there was actually a study, again by Linda Ganzini, who's written a lot about this, probably as much about this and what's going on in Oregon as anybody. She also claims to be neutral, she certainly is not coming from one camp or the other.

But the so-called Compassion and Choices folks gave her the right to interview 56 people who had been given massive overdoses by their doctors. And 25% of those people fit the criteria for major depressive disorder. 23% fit the criteria for anxiety disorder. Yet they were given prescriptions for overdoses.

Now many doctors are starting to say, just like Dr. Nitschke did yesterday with the three women who killed themselves, “Well, they'd been thinking about it for months, therefore it was rational.” And if it's rational, it's okay.

Well, I teach about depression to medical students, and residents and even practising doctors. And I'm good at it. I like to do it because it's so efficacious. It doesn't matter if a person is depressed because they had a heart attack, because they have a diagnosis of cancer, or they're like my son who was suicidal a couple months ago because he had had bad things happen in his life. And no matter which one of those people I help take care of, my help with their depression changes their thinking about the worth of their life.

Part of it is my skill in being able to detect it, ask appropriate questions, tell me what's going on, this must be really troubling, I want to hear about it. I become a doctor as drug, if you will. I know how to use the tools, whether it's behavioural therapy, cognitively, or it's the tools for anti-depressant.

And they work. It doesn't matter if you're depression's because of cancer like my wife, or because of other reasons and exigencies in your life, they work. I work with psychiatrists if I have to, with people who are really difficult to manage. There are all kinds of tools.

I'm not trying to get into that, I'm just simply saying all doctors don't have that interest, or that skill, and yet they're licenced to take people's lives, essentially practising vending machine medicine as Dr. Ganzini proved with her peer-reviewed publication in a psychiatry journal. It's on Medline, I've got the references here. You just click on the link. Hopefully, Mary will give you the handout so all these things, you can actually look and check on yourself.

And I would offer to you now, too, if you are in doubt about where to get these resources, so people can be informed with what are the facts rather than the hyperbole, the euphemisms, the fear mongering, then please contact our organisation or me personally directly, and I'll be glad to try to help you.
Only five percent of the doctors over the past 17 to 20 years have been referred to psychiatrists despite the fact that we've documented that 25% of them, based on Linda's study, are depressed. In some years, zero are referred. In fact, some doctors would say, well, of course they're depressed, they're terminal. Well, I would ask you, who in this room is not terminal?

Even the Supreme Court of the United States had that argument, what is the point here? If you're a diabetic and you stop taking your insulin, you're terminal. If you're somebody needing dialysis, and you decide not to use, which, by the way, these are your right, you're terminal. So you can have a juvenile diabetic who's terminal, simply by him or her not taking her prescribed medications.

**Myth Number Nine, it involves doctors who don't know the patient well.**

Often, this was pushed because it's important the doctor be involved because we can do it with real intimacy, because we know the person personally, we can prevent the abuses. Well, last year, one doctor did 25 deaths.

I can assure you that's not the statistical frequency of people who are terminal in your practise, even if you're a palliative care specialist. Let alone someone like me who's a family physician who sees all comers from conception to natural death.

The average length of time over the entire course of Oregon's experience is 13 weeks and falling. Because when you have people like Brittany Maynard come to Oregon as a suicide tourist, you don't have a lot of time to know who that patient is.

You don't have the ability to do what I can do with my patients when I know them for 18 years and they get in a funk and they're thinking about just not having even treatment, and they'd be dead in hours if they don't have treatment. And I can convince them that their life's worthwhile because I have a history with them and can reflect their worth even when they don't have their worth in their own minds.

Already, the ink is not dry as I said in California, and there's a drive-in clinic that's opened in San Francisco. It'll be $200 for the first visit, and if you qualify and he follows through with it, it'll be $1,800. There is the website of Dr. Lonny Shavelson, who believes he's doing a service to people with a drive-in death clinic.

You can't make this up. I mean, they say there's no slippery slope. You have to have your eyes closed, your head in the sand, your ears closed to say that. Again, even if there is a hard case that might bring your sympathy, aren't we also supposed to look at the societal impact of these measures?

So finally, I want you to understand that I believe there's a difference when people say these things and what they're really saying, so when patients say to me, "I want to die", tell me about that, because what they might be saying is "I feel useless".

Margaret mentioned the burden. "I don't want to be a burden." Are you a burden? And my response is, No, Mom, you're not a burden. This is literally my mother.

And I said, Mom, what's going on, what's wrong? I have hip pain. Well, Mom, did you take your medication? Yes. When? Yesterday. So I go and get her a pill, a simple, first-step pill before we get to the opioids. 20 minutes later, she says, Oh, I can't believe it, my hip doesn't hurt.

Because her mentation was not clear enough to know that she had to take it consistently to be free of pain. And then she starts feeling like, I might as well be dead, I'm a burden, why doesn't God take me? All of these messages which are the content expressed, but the real issue is I'm hurting, and the response to hurting is to try to help the hurt.
I don't want to be on a respirator like my patient Lonnie, (he said I could use his name). I said, you're not going to be trapped on a respirator, and I have to reassure him about his fear.

My wife was concerned about air hunger, because she had the spread of her cancer from the womb and the muscle in her womb to her lungs. I said, Marlene, we can get through this. We have drugs that help with the air hunger. And we did, we got through the air hunger. And the last night of her life was actually one of the most peaceful.

By the way, I had no idea she was going to die the next morning. Even when we woke and I helped her go to the bathroom, I did not know she was going to die in two hours. Again, I knew she was imminently going to die sometime in the next few weeks, month. But I didn't know it was that day, because doctors don't know. I didn't know, and I lived with her, I knew her.

I've lived a long life already, that's the completed life argument. Well, they're afraid you can't go on, no, we can do this, and you've got to encourage each other. I might as well be dead. Maybe you're saying nobody really cares about me.

So here's a person, and this is a real-life story of how does a doctor respond.

This doctor on the left is Dr. Ken Stevens. He was the head of the radiation oncology department for 40 years at Oregon Health and Science University. He's also a co-founder of Physicians for Compassionate Care.

Jeanette Hall is a patient who was diagnosed with terminal colon cancer, and she had voted for assisted suicide. It was Measure 16 in Oregon. She was not wanting treatment. She came to Dr. Stevens saying, I'm not here for treatment, chemotherapy, I'm just here for the pills. This is what she has to say.

[Jeanette Hall] I couldn't believe death was at my door. And it thought, surely that can't be right. Death, he's saying six months to a year to live. It was life and death. I just knew I didn't want to suffer. One day I was bleeding so bad that they had to take me by ambulance to the hospital. The
doctor came in and said, Jeanette, you have colon cancer, and it didn't hit me. I just went, I have to go, I have to take care of my mother, I have to do my work, I have to go. And they said, you don't understand. You have cancer.

[Dr. Ken Stevens] When I first met Miss Hall, she had been evaluated by her surgeon. Her surgeon had determined that the cancer was inoperable, meaning that it could not be cured with surgery. So he referred her to me for radiation and also chemotherapy. She immediately told me, she said, I'm not here for the treatment, Dr. Stevens, I'm here for the pills. Two years ago, I voted for Oregon's assisted suicide law, and that's what I want, I'm here for the pills. I don't want radiation, I don't want chemotherapy.

[Jeanette Hall] I reached that point without hope. I was sinking into depression.

[Dr. Ken Stevens] I said, Miss Hall, would you come back and see me next week, let's talk about this some more. I was able to get better acquainted with her, and I learned that she had a son who was attending the Police Academy, he was in his late 20s. And I said, Miss Hall, wouldn't you like to see him graduate? He was single, I said, wouldn't you like to see him get married? Does he know what you're considering here?

[Jeanette Hall] It made me realise, there was hope. There was something to live for. And I thank him for that one sentence that brought me back to reality and made me try.

[Dr. Ken Stevens] So she did accept the treatment. She received chemotherapy and radiation. She lost her hair temporarily, but the tumour just melted away.

[Jeanette Hall] Dr. Stevens, if you had believed in physician-assisted suicide, I wouldn't be here. You didn't give up on me, you didn't abandon me, and that's why I'm still here. These are the kind of doctors we need that give you hope when your hope is gone. Here I am, 15 years later, and I want to see my son's graduation. It's great to be alive.

So on the Mac (laptop screen showing end of video) it says, "I took the ancient oath to do no harm. I'll always care for my patients and never abandon them. "For me, helping them commit suicide would be the ultimate abandonment," Dr. Ken Stevens.

It's now 18 years for her.

All of us know patients who were told they only have three to six months, nine months, a year to live, and they were alive and kicking sometimes a year or two later, 17 years later, or like Jeanette Hall, 18 years later at this point. It's kind of funny when she actually talks about, he talked about, don't you want to see your son graduate, get married, and son quips back, well, I did graduate, but I haven't gotten married yet.

So Number 10, physician-assisted suicide is the solution to suffering. No, it's not.

If a person's in physical pain, treat the source of pain. If a person's lonely, provide companionship. That was what the original hospice, that's what Mother Teresa was about, that's what we should be about. If a person doesn't value their lives, work to reflect their inherent value. That's what Chochinov, a palliative care specialist in Canada, the Chochinov technique, espouses. It's in the peer-reviewed literature.

This is what we should do, follow Chochinov, not follow Nitschke.

If a person doesn't value their life, work to reflect their value just as we do everyone who's not labelled terminal, even though we're all terminal. And if a person's fearful, like my wife was about air hunger and her lungs not being able to breathe with the cancer that had spread there, reassure them and address those fears with the best technology we have today.
The solution to suffering is not to end the life of the sufferer. So just to review, assisted suicide is dispassionate care by doctors who often promote assisted suicide.

Now don't get me wrong, I'm not saying they want to be dispassionate, I'm just saying their behaviour ends up being that way. Whether you live or die is your choice. And you're essentially acting like the individual on an island.

It's dangerous, I've shared with you some of the problems. There are people who are standing to inherit wealth, caregivers who deposit $90,000 in their account a day or two after he died. She was indicted, she had other nefarious activities which is why she was found out, not because of this issue of the caretaking.

She took the house of the man and put it in her name and was going to sell it. This happened in Bend, Oregon. I can only share the ones that leak out because the government is doing such a lousy job of following these things, even though worldwide Oregon is reported as everything's going swimmingly.

There are numerous problems, I said.

There are no safeguards, as Margaret said at the outset, and if there are, they're like tissue paper and really like wet tissue paper. They really have no basis for being enforceable. And I believe there are compassionate approaches to hard cases and difficulty at the end of life. And even if I'm wrong and there's some hard case that is outside of my 40 years' experience, fair enough, but it's not worth the danger to society and the change in the doctor-patient relationship and trust in the doctor-patient relationship.

I thank you all for your time. I'll be glad to stay as long as you like to answer questions or hear your comments.

END