A Guide to the Debate on Voluntary Assisted Dying in Australia July 2021
Go Gentle Australia is an expert advisory and health promotion charity founded by Andrew Denton to spark a national conversation about end-of-life choices, including voluntary assisted dying. Our focus is on promoting evidence that is based on established facts and peer-reviewed research.

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Go Gentle Australia has been established to help relieve the distress, helplessness and suffering experienced by Australians with untreatable or terminal illnesses, their families and carers.

If you would like to learn more, visit us at gogentleaustralia.org.au
## Contents

**Introduction**
The Debate Ahead: Clock and Bomb  

**Part A: Some facts**
- Why VAD now? The unintended consequences of our existing laws  
- A historic level of consultation  
- Who this law is for: The limits of palliative care  
- Beyond pain  

**Part B: Questions raised by MPs**
- Will this law encourage elder abuse and coercion?  
- We should just resource palliative care better  
- VAD won’t lead to doctor shopping  
- Can we trust doctors to do the right thing?  
- Why VAD is not shown as the cause of death  
- What if there is a ‘wrongful death’?  
- Feeling like a burden is not the reason people choose a voluntary assisted death  
- Why a different consideration for people with a neurological disease?  
- Why a waiting period between first and last request?  
- The difference between VAD and suicide  
- VAD medication is reliable  
- Why a person is assessed on suffering and not just pain  
- How the Victorian VAD law works: The case study of Kerry Robertson  

**Part C: Four questions for legislators**
- Does this law make end-of-life practices safer?  
- Does this law protect religious freedom?  
- Does it make end-of-life practices more equitable?  
- Does this law meet community expectations?  

**Part D: Why facts matter in this debate**  

**End notes**
Introduction

The debate ahead
Clock and Bomb

We all know what a clock is. A precisely made piece of machinery, consisting of many parts, designed to do one thing: tell the time accurately.

The voluntary assisted dying bill you are about to consider is like a clock: A piece of legislation – designed with careful input from public health experts, senior clinicians and others – to offer terminally ill people in the last stages of their illness (and only those people) a choice about the manner and timing of their death.

It is very similar to the VAD ‘clock’ that has been operating effectively and uncontroversially in Oregon since 1997, and, since June 2019, in Victoria.

What happens if you take a clock apart and put all its component pieces on a bench?

It no longer looks like a clock. In fact, it becomes possible to argue that the pieces, when examined individually – this spring, that gear – may not be parts of a clock at all. They might be a bomb.

In the debates that lie ahead, opponents of this legislation – just as they have done in other parliaments – will use every tactic they know to try and make you forget that VAD is a clock, and to think of it, instead, as a bomb.

They know that, in their opposition, they represent only a small minority of Australians. They know that years of credible, peer-reviewed evidence from overseas shows that the ‘clock’ works as it is meant to. They know that the doctors and nurses you have heard from, who’ve talked about the need for this law and the protections it provides, are hard to deny. And they know that VAD is now legal in a majority of Australian states, the arguments against having been put under the greatest scrutiny and then rejected.

Faced with these realities, they have only one tactic. Use the parliamentary debate, which allows that the ‘clock’ of legislation be pulled apart and examined, to make you lose sight of the fact that it was ever a clock in the first place.

To do this, they will employ a debating technique you may have heard of, known as the ‘Gish gallop’.

The Gish gallop is a technique used during debating that focuses on overwhelming an opponent with as many arguments as possible, without regard for accuracy or strength of the arguments.¹

How does it work? Here’s New Hampshire legislator, Nancy Elliott, spelling it out to local campaigners at an international anti-euthanasia convention held in Adelaide in 2015:²

When you have lots of arguments, if one argument gets blown out of the water, you still have more, and each argument will reach somebody else... You only have to convince legislators that they don’t want this bill. You don’t have to win their hearts and minds. All you have to do is get them to say ‘not this bill’.

In the long hours of debate that lie ahead, through the blizzard of arguments that will confront you, the crucial thing to keep in mind is this:

Who is this law for, and how does it work to help them – and them specifically – at the end of life?

In other words, don’t lose sight of the clock.

– Go Gentle Australia.
Part A | Some facts

Why VAD now? The unintended consequences of our existing laws

After years of inquiry, the respective parliamentary select committees on End of Life Choices in Victoria, Western Australia and Queensland found that too many Australians are experiencing profound suffering as they die. Victoria’s parliament, acting on their committee’s recommendations, passed a Voluntary Assisted Dying Act in 2017. Western Australia, after a bruising parliamentary debate, followed suit in 2019. Since then, Tasmania and South Australia have also debated and passed their own VAD laws. But for other Australians living in states and territories without VAD laws, the choices currently offered them can be grim.

It is legal, if you are dying and suffering beyond medical help, to end your suffering by committing suicide. Most often this happens violently and alone. According to statistics from the state coroners, around one in ten suicides in any year is carried out by people with terminal or debilitating chronic illness. Each month in Queensland, for example, around seven terminally ill people take their own lives.³

It is legal to refuse all medical treatment, food and water, in the hope that, aided by starvation and dehydration, your disease will more quickly take its course. Your choice to do this will be medically supported. It can take days or weeks, and is not defined in law as suicide.

It is legal for a doctor to slowly drug you into a coma while your family waits for days, or weeks, for you to die. This can happen without your consent.

In other words, the law already recognises that a dying person has a right to hasten her or his own death in order to bring an end to suffering. Existing law means that such choices can be cruel, causing great trauma for the person dying, families, and carers.

In states without a VAD law in effect, it is not legal if you are dying and suffering beyond medical help, to end your suffering painlessly and quickly with the help of a doctor, at a time of your choosing and with an opportunity to farewell those you love.

As a patient, you have no legal right to insist that a doctor gives you more, or faster, pain relief. That decision is entirely up to the doctor whose personal beliefs you may not share.

Only in states that have passed assisted dying legislation does the law protect doctors who act compassionately to relieve intractable suffering by accelerating death.

In contrast, in states where there is no assisted dying law, the status quo means individuals who want to hasten their death must keep their plans to end their life secret from family to avoid them becoming an accessory. The law compels terminally or hopelessly ill people intent on taking their own lives to do so before they lose their physical or mental capacity.

THREE AUSTRALIAN PARLIAMENTARY SELECT COMMITTEES, WORKING INDEPENDENTLY, HAVE RECOMMENDED ASSISTED DYING BE MADE LEGAL

The length of life lost in such cases is a tragedy for the individual and all who loved them. This is why the three parliamentary committees looking at the issue recommended, by a majority, that VAD be made law:

• So that we have a legal right to be protected from unnecessary suffering at the end of life

• So that doctors can be protected in law, and guided by regulation, when doing what many currently do in the shadows

• So there can be more compassionate choices than our current laws allow if we are dying and suffering intolerably.
Over the last five years, the landscape in Australia around the debate on voluntary assisted dying has changed dramatically.

In this time we have witnessed three of the most comprehensive examinations of end-of-life care ever undertaken. Though conducted independently, these cross-party parliamentary inquiries, in Victoria, Western Australia and Queensland, found Australia’s existing end-of-life laws to be inadequate. All recommended, by a clear majority, that VAD be legalised as part of a broader approach to better end-of-life care, including more resources for palliative care.

The most recent of these select committee inquiries, in Queensland in 2019/20, received more than 5,000 submissions and heard from hundreds of people via public consultations. People of all ages and from all walks of life described with great clarity the damage being done in the absence of VAD laws.

In Western Australia, the successful Voluntary Assisted Dying Act (2019) was the result of a similar inquiry process – one of the biggest public and community consultations ever undertaken in WA. Running for more than 12 months, the 2018 Joint Select Committee Inquiry into End of Life Choices received more than 700 submissions and held 81 public hearings.

The Queensland and Western Australian processes, in turn, were built on the foundations of Victoria’s groundbreaking inquiry, held over 10 months in 2016. This committee considered 1074 submissions and heard from 154 witnesses. Members travelled to jurisdictions around the world where forms of voluntary assisted dying already existed to collect evidence.

• 3 PARLIAMENTARY INQUIRIES
• 200+ PUBLIC HEARINGS
• A RAFT OF INTERNATIONAL, PEER-REVIEWED EVIDENCE

The successful bills in WA and Victoria were further aided by the work of Ministerial Expert Panels. These panels, consisting of palliative care and other medical experts, former state and federal presidents of the AMA and expert advisors from the legal, disability and Indigenous communities, heard from hundreds of participants through public forums, roundtables and other meetings and received thousands of written submissions. In Tasmania, the Independent MLC who introduced the Bill consulted widely in Australia and overseas, and conducted scores of public forums and town hall meetings.

Supporting these forensic investigations and consultations is a raft of international, peer-reviewed evidence gathered from multiple jurisdictions with VAD laws in place. Jurisdictions where, after 10, 15 and even 20 years, there is still widespread public and political support for assisted dying; where fears about the abuse of the vulnerable have not been realised and where there is no loss of faith in doctors or degradation in palliative care – in fact the opposite.

Few, if any, laws tabled in Australian parliaments have been underpinned by such extensive consultation and expert thought.
Who this law is for
The limits of palliative care

While palliative care accommodates the needs of the majority of dying people, even the best-resourced service cannot relieve the extreme suffering some endure. This fact is acknowledged universally by responsible organisations and professionals involved in the palliative care industry. Western Australia’s Parliamentary Committee found that:

It is clear from the evidence that even with access to the best quality palliative care, not all suffering can be alleviated. Palliative care physicians themselves acknowledge this.4

In Palliative Care Australia’s own words

While pain and other symptoms can be helped, complete relief of suffering is not always possible, even with optimal palliative care.5

By its own calculations, the number of patients truly beyond palliative care’s help is about 4%.6 Other experts put this number higher at between 5 and 10%.7 The suffering, as recorded in evidence to the parliamentary inquiries, can be savage.

In 2016, 70% of respondents to an Australian Medical Association (AMA) survey agreed with the statement that:

Palliative care and medical treatment cannot adequately alleviate the suffering of some patients8.

Although Australia has the second-highest ranked palliative care system in the world9, these responses reflect the reality of modern medicine which can keep us alive longer, but which still has no cure for diseases like cancer and motor neurone disease.

“EVEN IF GOOD, MODERN PALLIATIVE CARE WAS AVAILABLE FOR EACH AND EVERY TERMINALLY ILL PATIENT – WE WOULD STILL HAVE THE ‘NIGHTMARES’.10

CLIVE DEVERALL
PALLIATIVE CARE WA FOUNDER
Beyond pain

Dr David Grube, a GP from Oregon with 35 years’ experience, who has worked within that US state’s Death with Dignity Act since its inception in 1997 and who visited Australia in 2019, explains from his experience some of the conditions that are difficult to palliate and which may lead a person to request VAD (see page 13).

Dr Roger Hunt, a specialist in palliative medicine for three decades, who sat on the WA Ministerial Expert Panel, explains how suffering is much more than just pain:

Dying people have varied and sometimes intense suffering, including physical, emotional and existential suffering. All surveys of palliative patients, including those in most renowned hospices, show they experience multiple concurrent symptoms. Severe refractory symptoms, including suffocation, pain, nausea and confusion, requiring palliation with deep sedation, have been reported in up to 50% of palliative care patients.

In his book, The Nature of Suffering and the Goals of Medicine (2004), Dr Eric Cassell, Emeritus Professor of Public Health at the Weill Medical College of Cornell University, a member of the Institute of Medicine of the National Academy of Sciences and a Master of the American College of Physicians, describes suffering as:

An affliction of the person, not the body.

Prof Cassell insists:

Not all suffering can be relieved, no matter how good the care ... and with even the very best treatment the suffering of some patients is terrible and unremitting. Experienced clinicians know that even if you think you have seen the worst suffering possible, given time you will see someone suffering even more. (Emphasis is ours)

Suffering at the end of life can be, for some, a condition that is ultimately impossible to palliate. Bone cancer, for instance, can be excruciating and almost impossible to relieve.

For a deeper understanding of suffering at the end of life, we urge you to watch the film The Broken Hearted (thebrokenhearted.org.au). It includes the story of Peta Quinlivan from Busselton, WA, whose husband, Russell, died in 2016 of lung cancer:

If people spent all this time with someone who was suffering, they might have some idea of ‘he died peacefully’. I don’t think so. Yes he might have been looking like he was peaceful at the end, anyone on that amount of pain killers might look peaceful at the end and anyone that is so exhausted from what they’ve been through might look peaceful at the end, but what about all the part leading up to it that wasn’t peaceful?

The palliative care response to intractable pain and suffering is terminal sedation, a medically controlled process that is utilised close to the end of life, commonly without the explicit consent of the patient.

In order to avoid accusations of intending to hasten death, the delivery of sedatives and analgesics is slowly titrated upwards, which means that control of pain and suffering may take some days to achieve. As nutrition is withheld, it can take up to two weeks for the patient to die.

Advocates of law reform believe it would be more humane to assist a patient with unbearable, irremediable suffering by the administration of drugs (upon request) that cause death in minutes rather than starving to death over several days or weeks.

70% of doctors agree palliative care cannot adequately alleviate all patients’ suffering
<table>
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<tr>
<th>CLASSIFICATION</th>
<th>DISEASE</th>
<th>SYMPTOMS</th>
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<tbody>
<tr>
<td>Neurological</td>
<td>Motor neurone diseases: Amyotrophic lateral sclerosis (ALS), Progressive bulbar palsy (PBP), Progressive muscular atrophy (PMA), Primary lateral sclerosis (PMA).</td>
<td>Gradual paralysis towards complete inability to move, then ultimately, to breathe: no treatment</td>
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<tr>
<td>Cancer</td>
<td>Stage IV malignancy: beyond treatment; incurable</td>
<td>The symptoms of cancer are more significant than the cause. Many of these occur at the same time. Pain: constant and irresolvable, multi-system Bone metastases: cancer in the bone is the most painful Distention: accumulation of body fluids (abdomen) Open sores: often get infected Weight loss: this causes all other symptoms to be exaggerated Odour: bodily discharges from orifices and skin that cannot be controlled Disfigurement Incontinence: bowel and/or bladder Fatigue: constant and unremitting Systemic inflammation: a persistent ‘flu-like’ feeling</td>
</tr>
<tr>
<td>Pulmonary</td>
<td>End stage emphysema</td>
<td>Unresolvable shortness of breath/smothering</td>
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<tr>
<td>Cardiac</td>
<td>Congestive heart failure</td>
<td>Smothering/intolerable swelling/angina</td>
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<tr>
<td>Gastrointestinal</td>
<td></td>
<td>(Can also be side effects of treatments) loss of appetite/diarrhea/pain</td>
</tr>
<tr>
<td>Genitourinary</td>
<td></td>
<td>Incontinence/vaginal discharge/bleeding/recurrent infections</td>
</tr>
<tr>
<td>Central nervous system</td>
<td></td>
<td>Uncontrollable seizures/intractable headache pain</td>
</tr>
<tr>
<td>Haematologic</td>
<td>Leukemia, etc</td>
<td>Continuous bleeding/untreatable infections</td>
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<tr>
<td>Dermatologic</td>
<td></td>
<td>Open sores/painful/dressing change problems/infections</td>
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The elder-abuse-and-coercion argument suggests that family members or carers could pressure or ‘steer’ an elderly patient to request assistance to die, for instance, so that they may access an inheritance sooner.

A VAD law makes such abuse less likely, not more, by bringing practices that are currently happening in secrecy into the light.

It encourages better practice. It brings regulation and scrutiny to a space that is currently unregulated. It offers the vulnerable the protections they currently do not have.

Proposed voluntary assisted dying laws come with strict eligibility criteria and strong safeguards to ensure they apply only to those for whom the law is written. To qualify, a person has to have an extreme medical condition which, in the assessment of two doctors, independent of each other, is clearly leading to her or his death and which can no longer be treated in a way that will meaningfully ease their suffering.

WA’s Ministerial Expert Panel found that:

… Where voluntary assisted dying is practised, systematic reviews have found that coercion is rare. Data from the Netherlands and Oregon show that ‘members of vulnerable groups are no more likely to receive assistance in dying … with the sole exception of people with AIDS’. The researchers concluded: ‘the available data … shows that people who died with a physician’s assistance were more likely to be members of groups enjoying comparative social, economic, educational, professional and other privileges’. Anecdotal reports suggest that coercion from relatives is more likely to take the form of an attempt to dissuade the person from voluntary assisted dying.15

On 9 August 2019, the Journal of the American Medical Association, reported on Medical Aid in Dying (MAID) laws in Oregon and Washington State:

Most patients who acquired lethal prescriptions had cancer or terminal illnesses that are difficult to palliate and lead to loss of autonomy, dignity, and quality of life. Concerns that MAID would unintentionally target socially disadvantaged patients have not materialized.16

Representatives of peak elderly and disability groups in Europe and North America also report no abuse of their members under these laws.17

Proposed VAD laws in Australia are designed with multiple safeguards to protect against the risk of abuse of the vulnerable. They mandate that doctors involved in assessing patients for eligibility be specifically trained to assess possible coercion and also capacity.

Compare this to existing end-of-life practices – terminal sedation and voluntary refusal of food and fluids – where there is little, or no, regulation or
oversight, no central record kept and no requirement for doctors to examine either capacity or the potential for coercion.

The proposed laws also involve a level of real-time scrutiny – with serious penalties for abuse – which are unique in end-of-life care. It would be far easier to steer or coerce patients into ending their life by overdosing on their medication, or by encouraging them to suicide (options available now), than it would be via these laws.

Elder and disability abuse is a serious problem. It is only made possible by abusers acting in what they believe to be secrecy.

On 28 July 2018, the then AMA WA President, Dr Omar Khorshid, admitted that doctors already routinely hasten death, telling journalists: "We (doctors) shorten patients' lives regularly if that is the right thing to do at the very end of life".18

Under existing law, when Australian doctors illegally assist a person to die, we do not know if that person has been subject to abuse or coercion.

We also do not know – and do not ask – if a dying person’s decision to refuse treatment, food and water has been made under pressure, subtle or otherwise.

A VAD law brings these practices that are currently happening in secrecy into the light.

It encourages better practice. It brings regulation and scrutiny to a space that is currently unregulated. It offers Australia’s vulnerable the protections they currently do not have.

It is appropriate to raise concerns about elder abuse and the abuse and coercion of the disabled. But it is morally wrong in doing so to raise hypothetical concerns about the abuse of vulnerable people as a reason not to address evidence of actual abuse happening right in front of us.

The Victorian, Western Australian and Queensland inquiries clearly show that vulnerable people are being mistreated and abused within our medical system because of existing laws.

**It is abuse** to force vulnerable persons to suffer a painful and drawn-out death against their wishes.

**It is coercion** to leave a dying person the choice of only further treatment (which has become intolerable) or suicide.

**It is not the prospect of VAD laws that heightens the risk of mistreatment and abuse, but rather their absence.**

I HAVE NOT SEEN - AND I HAVE BEEN LOOKING, BELIEVE ME, I HAVE SEEN NO INDICATION OF ANY TYPE OF COERCION.

BETTY KING, QC, CHAIR OF THE VICTORIAN VAD REVIEW BOARD.
We should just resource palliative care better

This is not, and never has been, an either/or debate. Assisted dying and palliative care both have the same aims: to alleviate suffering at the end of life; to help people die well, but also and just as importantly, to help them live well and with dignity in their last weeks and months.

The reality is that both palliative care and VAD are important. However, the view that ‘we should not consider VAD until palliative care is fully resourced’ is a classic Catch-22. Opponents know they can always argue that there are never enough palliative care services, no matter how much money is directed to them, because they can forever shift the hypothetical baseline of what constitutes ‘enough’.

Regardless of resources, there are clear limits to the effectiveness of palliative care (as discussed earlier).

There are also patients with chronic and progressive illnesses – such as advanced MS or motor neurone disease – who, because of the long-term nature of their suffering, do not sit within the palliative care system.

As well, some patients do not wish to die a ‘medicalised’ death, even one in palliative care. According to the Grattan Institute, 70% of Australians wish to die at home but only 14% do.19

Why should these patients and their suffering be set aside to reach a hypothetical goal of palliative care perfection, when we have the means to help them now?

Although both proponents and opponents of VAD strongly support increased resourcing of palliative care, it is important to remember that Australia already has the second-best palliative care system in the world, just behind the UK.20

Even so, the evidence shows that a voluntary assisted dying law will lead to better palliative care.

A 2018 report by Palliative Care Australia on the international situation found:

“There is no evidence that assisted dying has substituted for palliative care due to erosion of safeguards ... if anything, in jurisdictions where assisted dying is available, the palliative care sector has further advanced.”21

It found that physicians sought to improve their knowledge and understanding of end-of-life care support services and there was increased access and funding.

Since the parliamentary inquiries into end-of-life choices, the Victorian, WA and QLD governments have committed extra funding to support and enhance community-based palliative care services.

That palliative care and VAD go together is borne out in Oregon and in Victoria, where 90% of terminally ill patients who choose a doctor’s help to die are also enrolled in hospice or palliative care.

The proposed Australian laws are not intended to replace palliative care. They simply add one more end-of-life option alongside palliative care, when even the most heroic efforts of doctors and nurses can no longer help and when patients, exhausted by both their illness and its treatment, have suffered enough.

FEARS OF LOSS OF REPUTATION, TRUST AND FUNDING ARE QUITE UNFOUNDED. PALLIATIVE CARE STANDS TO GAIN IN RECOGNITION AND SUPPORT THROUGH THE INTRODUCTION OF VAD IN AUSTRALIA.22

PROF IAN MADDOCKS
THE ‘FATHER’ OF AUSTRALIAN PALLIATIVE CARE
VAD won’t lead to doctor shopping

An argument against VAD is that a patient seeking an assisted death could simply go ‘doctor shopping’ until they found one who agreed to help them.

The first and foremost eligibility requirement to access VAD is a terminal illness. The number of doctors visited cannot change this fact. Speaking practically, it would be near impossible for a seriously ill patient to go doctor shopping. Taking the Victorian and WA laws as a guide:

First, in both the WA and Victorian laws, when deciding the eligibility of a patient for VAD, doctors must consult a second practitioner. The first refers the patient to the second for a consulting assessment. Both doctors must report to the Review Board.

So, doctor shopping, if it is to take place at all, is restricted to the patient’s choice of the primary, coordinating practitioner. There are three possible scenarios where the patient could conceivably ‘shop around’:

- The chosen doctor declines or is ineligible to act as the coordinating practitioner (leaving the patient very little choice but to look for another doctor). Note that the Board will be aware of this, since the chosen doctor must inform it.
- Usually, the request and assessment process ends as a result of the assessment of the coordinating practitioner. Regardless of whether or not the patient informs a new choice of coordinating practitioner of this prior outcome, the Board will be aware of what is going on, since it will have the report of the previous coordinating practitioner.
- The second, consulting practitioner assesses the patient as ineligible. Doctor shopping at this point could not help but arouse suspicion, since the normal procedure would be for the coordinating practitioner to refer to another consulting practitioner.

Doctors may complain that all this reporting ties them up in knots, but the important point here is that it acts as a major constraint on the likelihood that a patient will abuse the system. In this respect it is quite unlike the classic ‘doctor shopping’ scenario, where the patient:

- is mobile and can readily move from suburb to suburb
- relies on the fact that neither the new doctor, nor the authorities, are aware in real time of what they are doing
- is able to present clinically with an illness that cannot be readily investigated or confirmed on the spot by the new doctor
- expresses urgency in the need to have the required restricted drug.

Second, we are talking about people at the end stage of a serious illness. Their capacity to keep looking around for a ‘correct diagnosis’ – which would already be apparent from their medical records anyway – would be extremely limited. This is not a ‘lifestyle choice’. No matter how many doctors they see, the bar never lowers. They have to meet all the eligibility criteria to qualify – criteria examined in real time and compared against the opinion of a second doctor by the VAD Review Board.

That doctor shopping doesn’t happen is borne out by the most recent statistics of the number of VAD prescriptions written in North America.

In 2020, in Oregon, a total of 142 physicians wrote prescriptions for 370 individuals. In 2018, in Washington, a total of 158 different physicians wrote prescriptions for 267 individuals. The medication was dispensed by 61 different pharmacies. In 2020, in California, a total of 262 unique physicians prescribed 677 terminally ill adult patients.

The AMA paints a picture of ‘euthanasia enthusiast doctors’ being encouraged under these laws. Based on the figures above, any such doctor acting exclusively as a VAD physician (if they exist) would struggle to make a living out of the handful of patients they would see each year.
Can we trust doctors to do the right thing?

Every day we trust doctors to perform procedures and provide medications that have the potential to end lives. We trust doctors to act well, even in such unregulated areas of end-of-life care as terminal sedation. It is difficult to understand the argument that providing doctors with more regulation will lead them to be less trustworthy.

A doctor’s mission statement doesn’t change under this law. Doctors are asked to act as they ordinarily do – diagnose, prescribe and care for their patients. Remember, too, figures from Oregon and Victoria show that around one in three patients prescribed VAD medication choose not to use it. Having been offered the option, they elect not to take it. Ultimately, it is the person who acts, not the doctor.

Why VAD is not shown as the cause of death

When somebody dying of cancer chooses to hasten death by refusing further treatment, food and fluids, the death certificate shows they died of cancer. When a terminal patient dies as a result of life support being turned off, the death certificate does not show ‘life support turned off’ as the cause of death.

The same principle applies with assisted dying.

In their submission to the WA Ministerial Expert Panel, National Disability Services explained it this way:

At the centre of the discussion is the balance between privacy and confidentiality of the person and the need to collect information about voluntary assisted dying. In some communities there may be significant impact on extended family of a person who has died through accessing voluntary assisted dying.26

The panel also received feedback that raised concerns about potential stigma if third parties (such as funeral directors) who have links to the person and their family or community were aware that a death had occurred through voluntary assisted dying:

The certification of death should match that concerning death itself; sanctity and respect. VAD death certification should be as respectful and confidential as possible to ensure it cannot be used for any religious or political agenda of any faith or secular group. – Submission by the Anglican Social Responsibilities Commission.
What if there is a 'wrongful death'?

By definition, the only people who can access a VAD law are at the end stages of a serious illness from which they are clearly already dying.

In the WA parliamentary inquiry process, Nick Goiran MLC made much of so-called wrongful deaths in his dissenting Minority Report arguing against a VAD law. In it he insisted that there was ‘peer-reviewed evidence of abuse’. However, these claims have been reviewed and found wanting.

Analysis shows the bulk of the evidence used in this report was not peer reviewed but instead came from abbreviated versions of official reports taken out of context, newspaper stories and anecdotes told by anti-euthanasia physicians.

A detailed investigation of 26 allegations of ‘wrongful deaths’ presented in the Minority Report shows that only six out of 26 cases (less than a quarter) related to patients who may qualify for an assisted death (under the legislation that was being debated in Western Australia).

Twenty cases in the Minority Report were entirely irrelevant, because the subjects fell outside the proposed criteria.

Of the six cases which may have been relevant to the Australian legislation:

- One concerned a woman who was only ‘terminally ill’ because she was reluctant to undergo chemotherapy. When her doctor did what any ethical physician would do – engage her in a discussion about her options – she relented and accepted treatment. She is still alive.
- Three recount incidents that occurred in the Northern Territory over 21 years ago under highly idiosyncratic circumstances.
- One concerned a patient being injected while in a coma – an irrelevant concern for a law which requires patients with capacity to self-administer a lethal drink or consent to doctor administration, under strictly prescribed circumstances.
- One has been rejected as ‘lacking veracity’ by the CEO of the healthcare provider in which the incident occurred.
- In four of the ‘wrongful deaths’ presented in the Minority Report, the subjects are, as far as can be ascertained, still alive.
- In two cases, the subjects were not approved for assisted deaths and died by their own hands.


The threat of ‘wrongful deaths’ is also not supported by multiple systemic, peer-reviewed studies from overseas, accepted by several Australian parliamentary committees.

In making it the argument, opponents of VAD claim that a regulated system with a level of real-time scrutiny of doctors is less safe than the current unregulated system where doctors admit to illegally assisting people to die.
Feeling like a burden is not the reason people choose a voluntary assisted death

It is true that one of the many ways in which a dying person can suffer towards the end of life is by feeling like they are a burden on those they love. It is not true to say that this is the reason they are given the legal right to end their suffering through voluntary assisted dying.

When Oregon data is examined in context, being a burden is not the only – or even the main – motivating factor for choosing assistance to die. In fact, it is well down the list, the chief reasons being loss of autonomy, dignity and enjoyment in life.

However, none of these factors are the reason these people have chosen and been given legal access to VAD. There is only one reason they are eligible – they are dying. The diseases they are dying of – mostly cancer, but also congestive heart failure and neurological diseases such as motor neurone disease – can all be seen on the yearly reports published by the Oregon Department of Health.

Since its inception in 1997, nobody – not a single person – in Oregon accessed voluntary assisted dying because they felt like a burden. They ticked ‘being a burden’ as just one of their end-of-life concerns because they were dying. The same criteria will apply elsewhere.

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<th>CHARACTERISTICS</th>
<th>2020</th>
<th>TOTAL</th>
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<tbody>
<tr>
<td>(N=245)</td>
<td>(N=1,905)</td>
<td></td>
</tr>
<tr>
<td>END OF LIFE CONCERNS</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Less able to engage in activities making life enjoyable (%)</td>
<td>231 (94.3)</td>
<td>1,713 (89.9)</td>
</tr>
<tr>
<td>Losing autonomy (%)</td>
<td>228 (93.1)</td>
<td>1,725 (90.6)</td>
</tr>
<tr>
<td>Loss of dignity (%)</td>
<td>176 (71.8)</td>
<td>1,308 (73.6)</td>
</tr>
<tr>
<td>Burden on family, friends/caregivers (%)</td>
<td>130 (53.1)</td>
<td>905 (47.5)</td>
</tr>
<tr>
<td>Losing control of bodily functions (%)</td>
<td>92 (37.6)</td>
<td>822 (43.1)</td>
</tr>
<tr>
<td>Inadequate pain control, or concern about it (%)</td>
<td>80 (32.7)</td>
<td>522 (27.4)</td>
</tr>
<tr>
<td>Financial implications of treatment (%)</td>
<td>15 (6.1)</td>
<td>86 (4.5)</td>
</tr>
</tbody>
</table>

SOURCE: OREGON DEATH WITH DIGNITY ACT, 2020 DATA SUMMARY
Why a different consideration for people with a neurological disease?

Medical and health practitioners commonly use the word ‘terminal’ to describe a situation when an illness or disease is expected to lead to a foreseeable or imminent death.

Legislation in North America requires that a person who has a terminal disease be able to access voluntary assisted dying. The most common terminal diagnosis for accessing voluntary assisted dying in these jurisdictions is cancer. The European jurisdictions of the Netherlands and Belgium do not require a person to have a terminal disease to access VAD.

In WA, a timeframe to death was put on the law because, in the Ministerial Expert Panel’s view:

* Assisted dying is intended to provide an option that can limit suffering at the end of life, not a way to end life for those who are not dying.

While a timeframe to death is a feature of the Victorian and Western Australian laws, there is no necessity to stipulate a timeframe if the legislation makes it clear the law is only for people at the end of their life. The spelled-out time periods were deemed appropriate in VIC and WA, but each legislature will draft a bill according to what is most appropriate for their situation.

A timeframe to expected death is referenced extensively in palliative care in Australia, often supported by the Surprise Question (a screening tool used to identify people nearing the end of life: “Would you be surprised if your patient was still alive in 12 months time?”). It is consistent with existing end-of-life and palliative care policy documents, such as the National Consensus Statement: Essential Elements for Safe High-Quality End of Life Care, as well as various state strategies.30

However, there may be good arguments not to include a timeframe, especially for people with degenerative chronic or neurological conditions. Often the progression of these diseases is unpredictable - decline can happen swiftly and with devastating consequences, even in those patients whose death was not predicted for many months. Sometimes, too, the suffering of an incurable and irreversible disease is greatest in those who are not immediately dying: rather, they are condemned to even greater suffering over a longer period.

For example, a person with advanced Multiple Sclerosis, or with motor neurone disease, may face many years of extreme pain, loss of autonomy, indignity and mental anguish.

People with terminal and debilitating chronic illnesses make up around 10% of suicides each year in Australia, as indicated by evidence to the VIC, WA and QLD inquiries. These violent and lonely deaths will continue in our community as long as there is no law to offer these people a peaceful choice.

Ironically, in allowing people a humane way in which to control their circumstances, many will choose to live longer, knowing they have an option should the worst happen. They can plan to farewell those they love and not leave behind the scars of an ugly suicide.

Lawrie Daniel’s last letter to his family illustrates the extent of the suffering experienced by people when they see no other option but to take matters into their own hands.

Lawrie died alone in January 2018, without the opportunity to say goodbye to his loved ones face to face. He wrote: “If we had a compassionate Voluntary Euthanasia process in this country, none of this would have to happen the way that it has. I hope you can forgive me, and that you and the children won’t see this as selfish, but as self-care, and self-compassion in a country where I had no alternative.”

Watch Lawrie’s Last Letter here: www.gogentleaustralia.org.au/lawrie_daniel
Why a waiting period between first and last request?

Some argue against a minimum waiting period on the basis that, whatever time period is landed on, it is never long enough.

The legislation recognises the need to guard against impulsive decisions by people experiencing extreme physical and emotional pain to ensure they are not accessing voluntary assisted dying without proper consideration.

Medical practitioners routinely assess whether medical treatment decisions are properly considered by their patients. Under these laws, two doctors, with training in assessing both capacity and the prospect of coercion, will do just that.

The primary (coordinating) medical practitioner is best placed to judge the enduring nature of a person's request in the context of the trajectory of the condition. The independent secondary (consulting) medical practitioner is best placed to act as a safeguard to ensure that the judgement of the coordinating doctor is reasonable. The legislation allows that these assessments, combined with the requirement that a patient be 'at the end-of-life', provide the necessary protection to ensure requests are properly considered, while also taking into account a person's condition and likely deterioration.

The time between each step in the process would, of itself, provide some period for reflection. The person would always have the protection of being able to withdraw from the process at any stage.

That a person's decision is enduring would be shown through several of the proposed process steps: the initial request, participation in the first assessment, the written declaration, participation in the second assessment, the request that triggers the provision of the prescription and the request for administration of the medication (if not self-administered).

In most jurisdictions there is some form of prescribed waiting period of anywhere between nine and 17 days.

For example, under the Dying With Dignity Act passed in Oregon in 1997, a person seeking assistance to die was required to make a verbal request, wait 15 days and then make a written request. They then had to wait an additional 48 hours before obtaining the prescription.

Subsequent research showed, however, that one in five people who made their initial request in Oregon died before they are able to complete the process.

In 2019, in an acknowledgment that the intention of the law was not to force eligible patients to suffer unnecessarily while they waited, Oregon lawmakers acted.

They amended the legislation (the only change in 25 years) so that doctors could make exceptions to the waiting periods if the patient was likely to die before completing them. A similar provision has been built into the Victorian and Western Australian laws.

That this is a compassionate and sensible provision was shown in a number of submissions to the WA Ministerial Expert Panel.

... perhaps with the ability to reduce times if the patient is undergoing excruciating pain that cannot be relieved, or death is imminent within that period.

– Submission by the Anglican Social Responsibilities Commission

Bethesda is of the view that it is reasonable to waive the waiting period if, in the opinion of two medical practitioners, the person will die before the waiting period has elapsed (as in Victoria).

– Submission by Bethesda Health Care
The difference between VAD and suicide

The distinction between suicide and a rational decision to end suffering was clearly understood by New York’s chief medical examiner, Charles Hirsch, when investigating the deaths of office workers who jumped from the Twin Towers on 9/11.

Faced with a terrible choice – a slow, agonising death by fire, or a quick death by jumping – many chose to jump. Seeing this as a rational choice to avoid needless suffering, Hirsch refused to classify their deaths as ‘suicides’.

Voluntary assisted dying is not a choice between life and death. It is the choice of a terminally ill person about the manner and timing of their death (which is imminent and inevitable) and the suffering that must be endured.

This difference was underlined by the WA parliamentary committee:

It is important not to conflate suicide with assisted dying. It is possible to distinguish temporary suicidal ideation from an enduring, considered and rational decision to end one’s life in the face of unbearable suffering.

The Lifeline position on the linking of assisted dying and suicide is:

Words can cause harm. Any linkage between euthanasia and suicide has the potential to cause harm.

We recommend that any public debate surrounding euthanasia refrains from making the link to suicide, as this can provoke suicidal ideation.

The reality is that the absence of VAD laws contributes to suicides. Key findings of the WA parliamentary committee were:

[Finding 33] The prohibition of a peaceful, assisted death has driven some terminally or chronically ill individuals to suicide using violent means.

[Finding 34] Some individuals who suicide under these circumstances are driven to take their lives early. All deaths under these circumstances are tragic and very often traumatic and distressing to family, friends and first-responders.

[Finding 35] It is impossible to quantify the number of people who attempt suicide and fail. However, there is evidence that many do and are left further debilitated.

Of the many compelling pieces of evidence to come from that inquiry was this from the state’s Coroner: That one in ten suicides in WA in any year are by people suffering with terminal or debilitating chronic diseases. Evidence in the Coroner’s report included the following:

- 199 cases where people who had been diagnosed with a terminal illness committed suicide
- There were one hundred (100) cases identified where the deceased was diagnosed with a terminal or debilitating condition and experienced a decline proximate to their death.
- From 2012 to 2017, there was an average of 41 deaths of relevance reported per calendar year.

ONE IN 10 SUICIDES IS BY A PERSON WITH A TERMINAL OR DEBILITATING ILLNESS

The Victorian and Queensland Parliamentary committees made similar findings: In Victoria, prior to the introduction of that state’s VAD law, the Coroner estimated around 10% of suicides were by people with chronic, debilitating or terminal illnesses. In Queensland, figures from the National Coronial Information Service showed seven people with terminal or debilitating conditions kill themselves every month, amounting to 168 deaths in 2016 and 2017.
VAD medication is reliable

The success rate of assisted dying medication is extraordinarily high. In Victoria, the Statewide Pharmacy responsible for preparing and delivering all VAD medications in that state reports a 100% success rate and no cases of significant complications with the medications in the program's first 18 months of operation.

In North America, the success rate is similarly high.

**Oregon**

In Oregon, of 1,905 deaths between 1998 and 2020:

- 33 people had difficulty ingesting (or regurgitated) the medication (1.7%)
- 3 people had seizures after taking the medication (0.16%)
- 8 people (0.42%) had regained consciousness after ingesting DWDA medications
- 16 people had other unspecified complications (0.84%).

Of 1,905 deaths, 60 people (3.1%) had some complication (not necessarily serious and not necessarily due to a fault with the drug) during or after ingesting the medication. This equates to 2.5 incidents per year over 23 years.

**Washington**

Between 2016 and 2018, of 540 deaths, complications were reported in 21 patients (4%). This equates to about seven patients per year over three years (the period covered in the 2018 report). In 2018, of 195 deaths, there were eight reported complications (4%).

The regurgitations mentioned are not failures. They are simply cases where patients had a slight negative reaction to the bitter taste of the medication or had been nauseous prior to taking the medication, or had some form of blockage. Nonetheless, they died peacefully.

In the few cases in Oregon where patients awakened, they may have ingested only a portion of the medication before they fell asleep, or they may have mixed the medications too far in advance and let it sit on a shelf, which caused a decrease in potency or crystallisation. In one case, more than 15 years ago, an inadequate dose for the patient's weight was prescribed.

No medical procedure is perfect. A success rate well in excess of 95% is extraordinarily high.

The WA Ministerial Expert Panel also found that:

> Despite the concerns raised in the consultation, the Panel noted there is no evidence of misuse with voluntary assisted dying medications elsewhere in the world.
Why a person is assessed on suffering and not just pain

Suffering is an intensely personal experience and is not confined to physical experience or pain. Many symptoms experienced by people suffering life-limiting illnesses are more difficult for the medical profession to assess and treat. Even with the best palliative care, both dying and incurably ill people face a range of symptoms that are difficult to manage. This is widely acknowledged, including by Palliative Care Australia and the Australian Medical Association.

While Australia’s palliative care is highly regarded, it can never be completely effective due to intractable symptoms which arise from, for example:

- cancer invasion of nerve-rich areas such as the abdominal cavity, chest cavity, spine, pelvis, or throat that leads to pain and, if in the pelvis, possible incontinence of urine and faeces. Cancer growth also leads to obstruction of swallowing, obstruction of the intestine, with vomiting and, ultimately, starvation
- paralysing diseases of nerves supplying the muscles of the chest and throat that cause gasping or choking to death (such as with motor neurone disease)
- mesothelioma (an asbestosis-related disease that is incurable), which produces severe chest pain with each breath, causing difficulty in breathing and feelings of suffocation
- difficult-to-treat neuralgia that causes experiences similar to electric shock, with stabbing, agonising and jolting pain.

This range of symptoms is indicative, not exhaustive, and reveals that suffering is not confined to the final stage of a terminal illness.

If you’ve had an accident, and a doctor wants to find out how much something is hurting, she or he is not going to tell you – but ask you. Only the patient can determine the level of pain and whether or not it is tolerable. However:

- Under the laws in WA and Victoria, two doctors also have to agree that what you say is intolerable suffering matches their reasonable expectation, based on your medical history and symptoms, of what intolerable would be
- It’s important to remember that, because you must have a terminal (and, by definition, terrible) disease to access VAD, it is highly likely that the first doctor you consult has already been treating you for that disease, knows your medical history, knows the trajectory of your disease, and is well-placed to consider a claim of intolerable suffering
- Any doctor who receives a request for VAD, even if she or he has not been treating you for that disease, is going to consult your medical records. Proof is needed – because all the records of a lawfully approved VAD request will go to the Review Board for examination – that you have the disease you say you have and that it has reached the point of intolerable suffering
- Even then, should they be convinced – both doctors, independently – that your suffering is intolerable, they still have to canvas all treatment options with you
- This is not a one-off consultation. Like all complex medical treatments, it is an ongoing conversation with your doctor.
The following account was written after an extensive interview by Go Gentle Australia with Kerry Robertson’s daughters. A version can also be read in The Age.

The daughters of the first Victorian to use the state’s voluntary assisted dying law say their mother’s death was “beautiful and peaceful”.

Kerry Robertson, 61, died in a nursing home in Bendigo on 15 July (2019) of metastatic breast cancer. She was the first person to receive a permit under Victoria’s Voluntary Assisted Dying Act (2017) and also the first to see the process through to its end.

Ms Robertson’s daughters Jacqui Hicks and Nicole Robertson, who were at her bedside, said: “It was a beautiful, positive experience. It was the empowered death that she wanted”.

“We were there with her; her favourite music was playing in the background and she was surrounded by love,” Jacqui said.

“Palliative care did its job as well as it could. But it had been a long battle. She was tired, the pain was intolerable and there was no quality of life left for her.”

The sisters said the experience had reinforced their belief that anyone who has a terminal diagnosis, is suffering and in intolerable pain, deserves the choice of a voluntary assisted death.

“It is the most compassionate, dignified and logical option for those suffering in the end stages of life,” said Nicole.
Timeline of events

KERRY’S CANCER HISTORY

- **October 2010** – Kerry is diagnosed with breast cancer. Has lumpectomy, lymph nodes removed, radiation and chemo.
- **March 2015** – Kerry requests scans to be completed which reveal that her cancer has metastasized in her bones. She begins radiation and bone regrowth injections.
- **December 2015** – the cancer spreads to Kerry’s lungs.
- **January 2016** – Kerry starts a long-term chemo plan. Scans show that there are tumours in her brain.
- **December 2018** – Kerry is hospitalised twice to manage her pain, but this provides only short periods of relief.
- **March 2019** – the cancer spreads to Kerry’s liver. Managing side-effects is affecting quality of life. Kerry decides to stop all treatments.
- **May 2019** – Kerry is admitted to hospice, struggling with pain, nausea and vomiting, and an inability to walk unaided. Her pain medications are adjusted again, with no relief.
- **June 2019** – Kerry moves into a nursing home, begins to decline rapidly.

VAD PROCESS

- **19 June** – initial verbal request to coordinating doctor and initial assessment are completed.
- **2 July** – second verbal request is made to consulting doctor and secondary assessment completed.
- **7 July** – third verbal request, written request and person of contact paperwork completed and submitted.
- **9 July** – permit is granted, script is sent via registered post.
- **12 July** – appointment is made with state-wide pharmacy to meet with Kerry.
- **15 July** – medication is dispensed; Kerry takes it the same day.
There is a clear body of evidence demonstrating that assisted dying already occurs, despite being unlawful. This includes research that examines doctors’ intentions when administering pain-relieving medication and whether the provision of this treatment always complies with the law.

As discussed elsewhere in this booklet, an act done with an intention to relieve pain is lawful (even if death is foreseen), but the same act done with an intention to kill is not lawful.

Despite this, some doctors who are treating terminally ill patients intend to kill when they administer pain-relieving medication, and so will be acting unlawfully. Further, the palliative practice of ‘terminal sedation’ – where a patient is kept under deep continuous sedation to manage pain, while artificial nutrition and hydration is withdrawn or withheld ultimately leading to death – can give rise to legal ambiguity and has sometimes been equated to ‘slow euthanasia’.

Two adverse consequences flow from the fact that unlawful practices occur.

The first is that as these practices are unlawful, they are unregulated. Regulation promotes good practice and, conversely, there are dangers inherent in unregulated practices, particularly where they lead to people’s lives being ended. For example, which patients are acceptable for doctors to assist to die? What practices are acceptable to achieve this purpose? Are doctors covertly making quality of life assessments that result in a decision to end a person’s life?

Legalisation and regulation of assisted dying allows for the creation of appropriate safeguards and oversight to ensure, for example, that a decision to end life is made only by a competent adult.

The second consequence of the existence of unlawful practices is that the ongoing occurrence of such practices in defiance of the law has the consequence of bringing the law into disrepute.
Medical professionals can refuse to participate in VAD due to moral or ethical convictions without sanction or criticism.

Many doctors also have strong religious beliefs that guide their approach to medical care. Australia’s Catholic palliative care services are guided by the Vatican’s Instructions for Health Care Workers:

*Pain in the final moments of life can take on spiritual significance for the sick person and... can be accepted as... ‘union with the redemptive sacrifice of Christ’... it is therefore necessary to give an evangelical meaning to death.*

A voluntary assisted dying law respects the right of any doctor or nurse to conscientiously object. In so doing, it respects religious freedom.

At the same time it protects a patient’s right not to have those beliefs imposed upon them.

However, while an individual’s right to conscientiously object is fundamental to a law whose first word is ‘voluntary’, it is imperative that the rights of the people for whom this law has been written – the terminally ill – are in no way compromised by a blanket allowance to institutions to do the same.

In determining what level of institutional conscientious objection is allowable, parliament must weigh up competing harms.

On the one hand, is the moral harm done to those who conscientiously object to VAD.

On the other, is the physical, emotional and psychological harm done to those who are dying and who are in too much pain, or too debilitated, to be moved – or who simply don’t have somewhere else to be moved to.

In allowing an absolute right for individuals to conscientiously object to participation in VAD, the law acknowledges their right to be protected from moral harm. Yet it goes further: acknowledging that, in certain circumstances, institutions may also seek to be protected from moral harm by arranging for the transfer of terminally ill patients seeking VAD as they undergo assessment and, ultimately, administration.

However, the law draws a line – adjudging that there is a greater harm in refusing access to doctors and medication to those seeking VAD, but who are unable to be transferred due to the nature of their illness, or the lack of an alternative facility.

In that instance, the law seeks to protect the person suffering from that greater harm.

It is a difficult balance to strike, but a necessary one.

Not to do so is to, effectively, create two classes of citizens: those who are capable of being transferred and, therefore, able to pursue their legal right to VAD; and those who are not.

The fundamental principal of any law is that it should apply equally to all citizens. To make VAD accessible to some but not to others, simply by virtue of the severity of their illness or of their postcode, is to jettison that principle.

The inherit unfairness in a law, would be further felt by extending it to those who live in aged care facilities and retirement villages.

Often, these are people’s permanent homes. They have all the things that are important to them there. They have a community. This is where they live. A law which says that – in order to protect the feelings of others – they must leave their home at the very time they are at their most vulnerable is both cruel and unfair.
generally able to do so because they possess some privilege. It may be privilege in terms of education or it may be in relation to contacts and connections they have within the medical or veterinary professions. It may be the wherewithal and financial means to travel to an assisted dying clinic in Switzerland.

The operation of the law cannot be justified if a privileged few are able to receive assistance to die, but others cannot.

There is also the imbalance that currently exists in law between patients and doctors. Under existing law, doctors have total power to decide how much pain relief is given to a dying patient, at what speed and when. Clive Deverall, founding president of Palliative Care WA, summed it up this way:

And every day, in different settings, terminally ill patients – most with advanced malignancies and uncontrolled symptoms – are terminally sedated. Excellent. But that is the Lotto Life: the patient has to be lucky enough to find a doctor willing to help. And there is little or no transparency and a legal risk to the doctor and/or nurse.46

That doctors hold strong personal beliefs that may influence their treatment decisions is not at issue. It is not suggested that they are wrong to do so, or that they should be required to act otherwise. Any proposed VAD law, rightly protects and respects a doctor’s right to act according to her or his conscience in providing end-of-life care.

What is at issue is an existing law that enables doctors to impose those beliefs on dying people in their care. People who, in many cases, do not share those beliefs. People who are suffering as they die and who make persistent and rational requests to end their suffering quickly.
Does this law meet community expectations?

Australians have long accepted that they should be free to make their own decisions about how they live and end their lives.

Consistent, reliable opinion polling over two decades reveals support for voluntary assisted dying to be in excess of 70%. In recent years, that support has risen to nearly 90%.47 This is irrespective of variations in the questions and terminology used.

In the most recent polls, support has remained stable at between 82% and 88%, according to Newspoll (2012),6 The Roy Morgan Snap Poll (2017), The West Australian (2018 and 2019),49,50 and ABC’s Vote Compass (2019).

This support also extends into the medical professions. A significant number of doctors, nurses and allied health professionals support patient choice at the end of life.

The Royal Australian College of General Practitioners, the Royal Australasian College of Physicians, the Australian and New Zealand College of Anaesthetists, the Australian Medical Students Association, and the Australian Nursing and Midwifery Federation, have all either backed a VAD law or adopted a neutral stance.

According to a 2016 survey conducted by the Australian Medical Association (AMA), a majority of members (51.6%) agreed assisted dying “can have a legitimate role in modern medical care”.

Yet the AMA, which represents not even 30% of doctors (and less than 25% in some states) continues to oppose assisted dying, ignoring the views of a significant proportion of its membership.

In WA, two former state AMA presidents served on the Ministerial Expert Panel that provided recommendations on the VAD bill to the government.

Both have publicly distanced themselves from the official AMA stance, with one former president, Dr Scott Blackwell, declaring that he is disappointed with the AMA’s “alarmist and concerning” commentary.

[The commentary is] unnecessary and probably unrepresentative of the body of people who call themselves AMA members, and certainly of the body of people who call themselves doctors in WA.51

51.6% OF AMA MEMBERS AGREE THAT ‘EUTHANASIA CAN FORM A LEGITIMATE PART OF CARE’52
Ben White, Professor in the Australian Centre for Health Law Research at QUT, an independent researcher who has been working in this field for 18 years, has this to say about the assisted dying debate:

Voluntary assisted dying is a critical social issue. Both sides of the argument have a duty and responsibility to be transparent in their views and also to be clear about the evidence they are relying on. Where there is trustworthy, reliable evidence that sheds light on how voluntary assisted dying regimes work, it’s absolutely critical that law makers, parliamentarians, parliamentary committees, politicians, media, and policy advisers must have access to that, engage with that and understand how it can help make evidence-based law.

Some evidence will be more reliable than others and there is established ways to test this. Evidence which draws on a large body of peer reviewed [material]...

We can have more confidence in evidence like that than that which might be a handful of cases which are anecdotally reported.55

So, if you are looking for information and want to know if it is credible or not, a starting point would be to look at who is the author. If it is a government department, for example, you might have confidence in the information that is there.

Similarly, there have now been several parliamentary committee reports in Australia. All are available online and detail the evidence about assisted dying here and overseas. These are trustworthy sources of information.

A report55 prepared in 2018 by the conservative Institute of Public Affairs (IPA) and the progressive Per Capita think-tank, investigated 20 policy decisions in recent years by the federal Coalition government, the NSW Liberal government, and Queensland and Victorian Labor governments.
Despite differing outlooks, the IPA and Per Capita agreed on what were good and bad policy processes. At the very top of their list of good policy was Victoria’s voluntary assisted dying legislation. If this exercise were repeated today, Western Australia’s VAD law would similarly appear at the top of the list.

The director of policy at the IPA, Simon Brehny, said that in a period of “declining trust in politicians, democracy and institutions, it is essential now more ever that policy making is undertaken in a thorough and consultative manner.”

Good policy process – from actually undertaking cost-benefit analysis to having a detailed plan for how a policy will be rolled out – is not a left-right issue; it is an issue of competency.

According to Emma Dawson, the executive director of Per Capita Australia,

“Our assessment of the selected policies against the agreed criteria revealed the importance of taking a rigorous and consultative approach to policy development and implementation at all levels of government.”

We all have an extraordinary responsibility in debates about voluntary assisted dying and should:

- Look carefully at the evidence at the top of the evidence pyramid. (see page 28)
- Keep in mind the vast experience of the clinicians and others, who have informed the writing of these laws;
- Remember for whom the law is intended – those who are dying and suffering and who are seeking a choice about how cruel their death needs to be;
- Finally, never forget that this law is a clock.
End notes

11. Dr David Grube MD, visit to Western Australia, August 2019, facilitated by Go Gentle Australia.
14. Ibid. p. 289
22. Ian Maddocks, Correspondence with Go Gentle Australia. 2017
30. WA End-of-Life and Palliative Care Strategy 2018-2028 https://www2.health.wa.gov.au/~/media/Files/Corporate/general%20documents/Health%20Networks/Palliative%20care/WA%20End-
Go Gentle Australia Limited is a registered charity.

Go Gentle Australia is a health promotion charity founded by Andrew Denton. It was established to help relieve the distress, helplessness and suffering experienced by Australians with terminal illnesses, their families and carers. We are about a better conversation around death, dying and end-of-life choices, including voluntary assisted dying.

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