



Your rights when using health or disability services



Before you start



This is a long document.



It can be hard to read a document this long.



Some things you can do to make it easier are:



- read it a few pages at a time
- set aside some quiet time to look at it
- have someone read it with you to support you to understand it.

What you will find in here

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What is this document about?



This Easy Read document is from the **Health and Disability** Commissioner.



The Health and Disability Commissioner is also called the HDC.



HDC looks after the rights of people who use:

- Your rights when you use a health or disability service
- HX



- You have the right to:

- - have good care and support that fits your needs be told things in a way you understand
- be told everything you need to know about your care and support

Talk to the person you are not happy with

- Call 0800 55 50 50 and ask for a Health and Disability Advocate or email advocacy@advocacy.org.nz Call 0800 11 22 33 or email hdc@hdc.org.nz to make a complaint with HDC
- HDC protects the rights people have under the Code of Health and **Disability Services Consumers'** Rights.

health services

Protecting your rights



Everyone who uses a health or disability service has their rights protected under New Zealand law.



Every **provider** of a health or disability service has things they must do to make sure their service **respects your rights**.



A **provider** is a person or organisation that offers health or disability services like:



doctors



dentists



- counsellors
- medical centres.









To **respect** you means that someone else:

- treats you fairly
- is kind to you
- knows your rights
- does not ignore your rights.

Providers need to:

- tell you about your rights
- let you use your rights.

About your rights



You have 10 rights under
New Zealand law when using a
health or disability service.



This part of the document will explain what your rights are.



Right 1: Respect



Providers need to:

- treat you with respect
- make sure that you have privacy.



Privacy means not sharing information about you with other people without your permission.



Providers need to consider your:



social needs like friendships



- religious needs like going to church
- cultural needs like being able to talk to someone in your language
- ethnic needs like traditions.







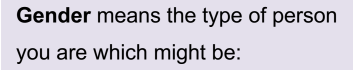


Right 2: Fair treatment

How well your treatment is done should not change based on:

- your age
- your **gender**
- your race
- if you are working or not
- if you are in a relationship
- who is in your family.

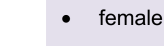




- the same as what you were called at birth
- different from what you were called at birth.

There are many genders including:













- nonbinary
- gender fluid.







How well your treatment is done should also not change based on:

- your beliefs
- your disabilities
- your **sexual orientation**.





Sexual orientation can mean:

- gay / lesbian
- straight / heterosexual
- bisexual / pansexual
- asexual.













You should be able to use services without being:

- made to do something you do not want to do
- threatened like someone saying they will hurt you if you do not do something
- harassed.

Someone **harassing** you is when someone keeps doing something hurtful when you have asked them to stop.

Harassment can happen in lots of ways like:

- calling someone names
- hitting someone
- sending mean emails.

Right 3: Dignity and independence



Services providers must respect your:

- dignity
- independence.



Dignity means the respect you have for yourself.



Independence means making choices for yourself.









Right 4: Appropriate standards

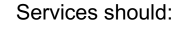
Appropriate standards means that services should be done:

- with a good quality of care
- by someone who knows what they are doing
- in a way that supports you to live the best life you can
- in a way that makes sure any treatment that might be painful is not any more painful than it should be.





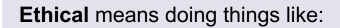




- follow the law
- make sure their staff do their jobs well
- make sure they offer good care
- be **ethical**.







- being honest
- making choices that are fair
- treating people well.



Right 5: Effective communication



You should be given information in a way that you can understand.



You should be able to ask for an **interpreter** if:



- you need one
- it is possible to get one.



An **interpreter** is someone who can explain what someone else is saying in a language you understand.







- held in a safe place
- done in a way that:
 - o is easy to understand
 - o works well for everyone
 - o is honest.





Communication being **honest** means:

- telling the truth
- telling someone why you think something
- not keeping information from people when they need to know that information.

Right 6: Information









You should always be told:

- about any health conditions you have
- how long a provider thinks it might be before you can use a service
- any information you might need to make a choice
- what providers find out as part of any:
 - o tests they do
 - o treatments you have.

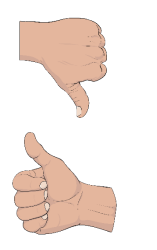


You should also be told if you might be part of any:

- teaching
- research.



Research is when people find out a lot about something.



You should be told everything about the choices you can make like any:

- things that could go wrong
- good things that could happen
- money you might have to pay









Side effects are things like a headache or feeling sick because of medicine or treatment.

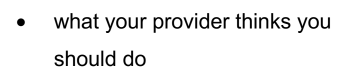


You should be given answers to any questions you have about services like:



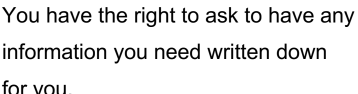
who the provider of the service is

what training the provider has had





how to ask what a different provider thinks



what has been found out because

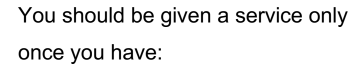


for you.

of research you were part of.

Right 7: Choice and consent





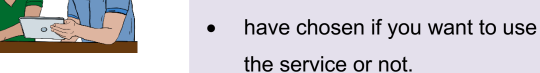


- made an informed choice
- given your **informed consent**.



An **informed choice** means that you:

- have been told any information you need or want to know about the service
- understand what this information means











- have enough information to make a decision
- agree to use the service
- understand what:
 - o using the service means
 - could happen if you use the service.



Providers should think that you are **competent** unless they have a good reason not to.











Competent means that you can:

- understand information
- think about what choice is best for you
- ask questions about things you are not sure about.

Some people are not competent to make informed choices.

If you are not competent to make an informed choice then you should still be allowed to:

- make the choices that you understand
- choose to give consent to things you understand.



Sometimes people cannot give their consent when it is needed like if they are unconscious.





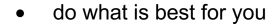
Unconscious means that you are not:

- aware of what is happening around you
- able to communicate / talk.

Being unconscious is like being in a very deep sleep that is hard to wake up from.



If a service is needed when you cannot give your consent then providers need to:





- try and find out if you would usually consent to the service
- talk with people who know you to find out what you might think.



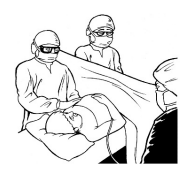
You will need to consent in writing if you agree to:

- be part of some research
- use a general anaesthetic









A **general anaesthetic** is a medicine that is used to make you:

- sleep during an operation
- not remember the operation happening
- not feel any pain during the operation.



An **experimental procedure** is a treatment that is still being tested.



If something might have very bad side effects then you need to say in writing that you understand what these side effects are.









You have the right to:

- not agree to a service
- change your mind about using a service at any time
- ask to change providers if it is possible
- make a choice about most future treatments.

Some treatments cannot have choices made about them before they are needed.







You have the right to choose:

- what happens to any parts of your body that might be removed as part of a treatment
- what happens to your bodily substances like your blood
- if any of your body parts can be:
 - o used
 - o stored.

Right 8: Support



You may bring a support person with you as long as it:





 does not go against the rights of other people.

Right 9: Teaching and research



You also have all these rights if you are part of:



- teaching
- research.

Right 10: Complaints taken seriously



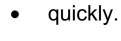
You can make a complaint about a provider in a way that works for you.



Providers must try to work out your complaint with you:



fairly





You should be told how making a complaint to your provider works.



Your provider should tell you:

- what is happening as they work on your complaint
- what happens because of your complaint.



HEALTH & DISABILITY COMMISSIONER

TE TOIHAU HAUORA, HAUĀTANGA

You should be told about who can support you with your complaint like:

- health advocates
- disability advocates
- the HDC.



You can find the full Code of Rights on the HDC website at:

www.hdc.org.nz/your-rights



Sometimes providers might not be able to meet all your rights.



Providers must always do their best for you even if they cannot meet all your rights.

Assisted dying



Some rights work differently for assisted dying services.



Assisted dying is end of life care that supports people who would like medical support to die when they choose.

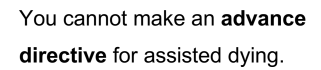


Under Right 6: Information means a health practitioner can tell you about assisted dying **only** if you ask them.



You will need to show that you are competent to make an informed choice about assisted dying.







An **advance directive** is a plan you write to tell your doctor what services you want in the future.



An advance directive is used if you cannot tell your provider what you want.



Your doctor does not have to perform assisted dying services for you.



Your doctor must still make sure you are cared for if they do not perform assisted dying services.



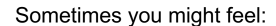
You can find out more about your rights and assisted dying on the HDC website at:

https://tinyurl.com/yr35nk6w

If you think your rights have not been met







- unhappy with a service
- that your rights have not been met.



To make sure that the problem is fixed quickly you need to first:

 talk to the person who provided the service





talk to the person in charge of the service.



You can ask other people for support in talking to the provider like:

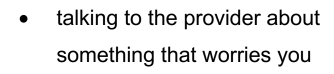
- your whānau / family
- friends
- an advocate from the Nationwide Health & Disability Advocacy Service.



You can find out more about the Nationwide Health & Disability Advocacy Service on **page 44**.



You should not have anything bad happen from:





• making a complaint.



If you feel that you are still not getting your rights met by a health or disability service you can make a complaint to the HDC.



A complaint sent to the HDC is dealt with in a way that:

- is fair
- works well.



HDC will work on your complaint as quickly as possible.





You can find an Easy Read guide to making a complaint in the document:

Form to make a complaint about your health or disability care

You can find this document on the HDC website at:

www.hdc.org.nz/making-a-complaint/make-a-complaint-to-hdc

How to contact us



You can contact the Health and Disability Commissioner by:



• phone: **0800 11 22 33**



• email: hdc@hdc.org.nz



We are open:





- Monday to Friday
- 8 o'clock in the morning to6 o'clock in the evening.

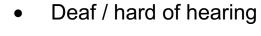


You can find out more about the Health and Disability Commissioner on our website at:

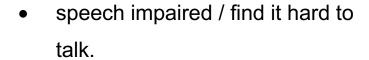
www.hdc.org.nz



If you find it hard to use the phone the **New Zealand Relay** service is for people who are:









You can find out more about the New Zealand Relay service at:



www.nzrelay.co.nz



You can also use Seeflow to record a video to send to us.



To use Seeflow go to their website at:

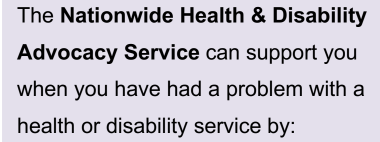
https://www.seeflow.co.nz/direct

How to contact an advocacy service



If you need more support to fix a problem you have had with a health or disability service you can contact the Nationwide Health & Disability Advocacy Service.







 talking with you about what you can do



- telling you about your rights
- answering your questions
- supporting you in making a complaint.













You can contact the Nationwide Health & Disability Advocacy Service by:

• phone: **0800 555 050**

• email:

advocacy@advocacy.org.nz

They are open:

Monday to Friday

8 o'clock in the morning to6 o'clock in the evening.

You can find out more about them on their website at:

www.advocacy.org.nz



This information has been written by the Health and Disability Commissioner.



It has been translated into Easy Read by the Make it Easy Kia Māmā Mai service of People First New Zealand Ngā Tāngata Tuatahi.



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