

Applying to the NDIS

Quick summary: If you want to become an NDIS participant you'll need to apply to the NDIS. There are some requirements you need to meet to be eligible for the NDIS. First, you need to be younger than 65 when you apply, and be an Australian citizen, permanent resident, or hold a Special Category visa that is a protected special category visa. You also need to live in Australia. Then, you'll need to meet the requirements for disability or early intervention.

You may be eligible under the disability requirements if you have one or more impairments that are or are likely to be permanent, and this substantially impacts your ability to do daily life activities. Your impairment must also affect your social life, or your ability to work and study. And, you must be likely to need NDIS supports for your lifetime.¹ [NDIS supports](#) are the services, items and equipment that can be funded by the NDIS.

Or you may be eligible under the early intervention requirements if you have one or more impairments that are or are likely to be permanent, or you're a child with developmental delay, and supports are likely to benefit you by reducing your need for supports in the future. We will also consider if these supports are NDIS supports.

Or you may be eligible under both the disability and early intervention requirements.

If you think you might be eligible, we can help you apply to the NDIS. We'll talk to you about your needs, current situation and what's important to you. We'll look at the information you give us to decide if you're eligible. If you're eligible for the NDIS, you'll become a participant, and we'll work with you to start [creating your plan](#). If you're not eligible, an early childhood partner or local area coordinator can help you explore and access government and community services. When we work with children younger than 9 and their families, we call this [early connections](#). When we work with people aged 9 to 64, we call this [community connections](#).

When we say 'you', we mean a person applying to the NDIS.

When we say 'we', we mean the NDIA.

When we say 'participant', we mean an NDIS participant who has met the NDIS eligibility requirements. Participants get funding in their NDIS plan to buy NDIS supports.

What's in this guidance?

This guidance covers:

- [What do we mean by applying to the NDIS?](#)

- [Do you meet the age requirements?](#)
- [Do you meet the residence requirements?](#)
- [Do you meet the disability requirements?](#)
- [Do you need early intervention?](#)
- [What about children younger than 6 with developmental delay?](#)
- [How do you apply to the NDIS?](#)
- [How do we decide if you're eligible?](#)
- [What happens after we decide?](#)
- [List A: Conditions that are likely to meet the disability requirements](#)
- [List B: Conditions that are likely to result in a permanent impairment](#)
- [List C: What if you're receiving disability support in Western Australia?](#)
- [List D: Permanent impairment/Early intervention, under 7 years. No further assessment required](#)
- [When do we make priority eligibility decisions?](#)
- [How do we weigh evidence of disability?](#)

What do we mean by applying to the NDIS?

Applying to the NDIS means doing things to find out if you're eligible to become a participant and start getting funded NDIS supports. There's a process to follow when you apply to the NDIS. As part of this process, we'll need to confirm your identity. Learn more about [how to apply to the NDIS](#).

For children younger than 9, we encourage parents and carers to talk to an [early childhood partner](#) about their child's development. They can help families connect to the right supports and let parents and carers know if applying to the NDIS is right for their child. Learn more about our [early childhood approach](#) and [early connections](#).

After you apply, we'll look at the information you give us and decide if you're eligible.

If we decide you're eligible, you'll become an NDIS participant. We'll then work with you to create your first plan, which will include funding for NDIS supports.

If you've been assessed as not eligible, there may be other supports available to you. For example, other government and community services. We can help you connect to these. If you're aged 9 to 64, check out [community connections](#). For children younger than 9 and their families, check out [early connections](#).

Are you eligible for the NDIS?

To be eligible for the NDIS, you first need to meet the [age requirements](#) and [residence requirements](#). This means you need to be younger than 65 when you apply, live in Australia, and be an Australian citizen, permanent resident or hold a Special Category visa that is a protected special category visa. You also need to live in Australia.²

You'll need to meet the requirements for disability³ or early intervention⁴. We may decide that you meet both the disability and early intervention requirements.

Under the law for the NDIS, we check if you're eligible based on your impairments, not your type of disability or diagnosis. When we say impairment, we mean a loss of, or damage to your body's function and/or structure. There are different types of impairments, including intellectual, cognitive, neurological, sensory and physical impairments, as well as impairments caused by a psychosocial disability.

To meet the [disability requirements](#), we must have evidence of **all** of the following:

- You have a disability caused by one or more impairments. This means your disability and impairments are linked.
- Your impairment is or is likely to be, permanent.
- Your impairment substantially reduces your functional capacity in relation to one or more daily life activities. These activities are communicating, socialising, learning, moving around, undertaking self-care, or self-management tasks.
- Your impairment affects your ability to work, study or take part in social life.
- You'll likely need NDIS supports for your lifetime.

To meet the [early intervention requirements](#), we must have evidence of **all** of the following:

- You have an impairment that is or is likely to be, permanent, or you're a [child younger than 6 with developmental delay](#).
- Early intervention will benefit you by reducing your need for supports in the future.
- Early intervention will benefit you by either reducing the impact your impairment has on your functional capacity or support your informal supports to build their skills to help you. Or help prevent the deterioration of your functional capacity or improve it.
- The early intervention supports you need are NDIS supports.

We'll use information in your application to [decide if you're eligible for the NDIS](#).

If you're eligible, you become an NDIS participant.⁵ The length of time you'll be a participant depends on the eligibility requirements you met, your individual circumstances and NDIS support needs. Learn more about [leaving the NDIS](#).

For general information about who's eligible, go to the [Applying to the NDIS factsheet](#). Check out our website for information about [how to check eligibility](#) and [children younger than 9](#).

If you have a psychosocial disability, find out more about [what information we need if you apply](#) under **Psychosocial disability**.

A psychosocial disability isn't a mental health condition or diagnosis. A psychosocial disability means you have reduced capacity to undertake tasks and activities of daily living due to your mental health.

Do you meet the age requirements?

To be eligible for the NDIS, you must first meet the age requirements. This means [you're younger than 65 on the day you apply](#).

How old are you?

You must be younger than 65 on the day you make your NDIS application.⁶

This means your completed application needs to be submitted to us before you turn 65. Learn more about the [application process](#).

If you're turning 65 soon and want to apply, [contact us](#) so we can help you apply on time.

Do you meet the residence requirements?

To be eligible for the NDIS you must then meet the residence requirements. This means we must have evidence of **all** the following:

- [You are an Australian citizen or permanent resident](#).
- [You live in Australia](#).

Are you an Australian citizen or permanent resident?

You must be an Australian citizen,⁷ or have one of two visa types that mean you can live here. A:

- [permanent residency visa](#)⁸
- [Special Category visa](#)⁹ that is a protected special category visa – this is only for some citizens of New Zealand.

Do you live in Australia?

You must live in Australia.¹⁰ This means Australia is your home and you spend most of your time here.

To help us decide, you need to give us evidence to show us you live here. If you give us consent to use your Centrelink record, that usually gives us enough evidence to decide if you live in Australia.

If you don't give us consent, you need to give us enough information to help us decide that you live in Australia. You'll need to answer these questions:

- [Where do you live?](#)
- [Where is your family?](#)
- [Do you work in Australia?](#)
- [Do you own property in Australia?](#)
- [How much time do you spend outside Australia?](#)

Some of these questions might not apply to you, but we must think about them when we consider whether you live in Australia.¹¹

We may also ask you other questions to determine whether you live in Australia.¹² For example, your family might be deployed overseas in the Defence Force, meaning you need to leave Australia for a while. In these situations, we'll ask you for more information about why you're leaving and can't return.

If you apply, we look at your whole situation when we decide if there's enough evidence to show that you live here. This will be a simple decision for us in most situations. But sometimes we may need to look at the questions below.

Where do you live?

We think about where you live, and your living situation.¹³ We'll look at whether you have more permanent accommodation in Australia than any other country.

For example, you might own a home or have a formal rental agreement in Australia. This is a good sign to us that you live in Australia.

Where is your family?

We also think about where your immediate family lives.¹⁴

We look at where you spend most of your time with them, face-to-face. We don't look at how you connect with your family on the phone or internet.

Do you work in Australia?

If you work, we look at where you normally work or make money.¹⁵ If you work or make money in Australia, that's a good sign you live here. If you don't work or are unemployed, we'll consider the other questions to determine whether you live in Australia.

Do you own property in Australia?

We look at what assets or property you own in Australia.¹⁶ We also see if you have an Australian bank account.

If you own assets or property here, it doesn't always mean you live in Australia. Your assets or property will need to show you have an ongoing connection to Australia. You don't live in Australia just because you own assets or property here.

How much time do you spend outside Australia?

We look at how often you go overseas, and how long you're outside Australia when you travel.¹⁷ We also look at why you travel overseas, such as for work, holiday or to live with family.

This is usually the most important point to help us decide if you live in Australia. You need to show us that you have a long-term and meaningful connection to living in Australia.

You don't need to stay in Australia all the time. You can still go on holiday or sometimes work overseas.

You'll need to show a stronger connection to Australia than other countries if you spend a lot of time overseas.

If we decide you're eligible and create your plan, there may be times you can't use your NDIS funding overseas. This is usually after you're overseas for more than 6 weeks unless we give you more time. Learn more about when you can't use your plan in [Our Guideline - Your plan](#).

Do you meet the disability requirements?

You meet the disability requirements if we have evidence of all the following:¹⁸

- [Your disability is caused by an impairment](#).
- [Your impairment is or is likely to be permanent](#).
- [Your permanent impairment substantially reduces your functional capacity](#) in relation to one or more of the following activities as a whole: communicating, socialising, learning, moving around, undertaking self-care, or self-management tasks.

- [Your permanent impairment affects your ability to work, study or take part in social life.](#)
- [You'll likely need NDIS support for your lifetime.](#)

If you give us evidence you have been diagnosed with one or more conditions on [List A](#), we'll likely decide you meet the disability requirements.

If you meet the disability requirements, it's likely you'll need NDIS supports for your lifetime. This means you won't have to prove your disability every time we reassess your plan.

If at any time your NDIS support needs or situation changes, we may need to check your NDIS eligibility. We'll talk with you if this happens.

Learn more about how we check [if you're still eligible for the NDIS](#).

Is your disability related to an impairment?

When we think about your disability, we look at whether any reduction or loss in your ability to do things is because of an impairment.

An impairment is a loss of or damage to the body's structure and/or how the body functions. We'll look at:

- your body's functions
- your body's structure
- how you think and learn.

It helps us understand how your disability impacts your everyday life.

To meet the disability requirements, we must have evidence your impairment relates to at least one of the following impairment categories:¹⁹

- **intellectual**
- **cognitive**
- **neurological**
- **sensory**
- **physical**

You may also be eligible for the NDIS if you have a **psychosocial disability**.²⁰ This means you have reduced capacity to do daily activities and tasks due to your mental health. Your psychosocial disability might vary at different times in how much it impacts your daily life. Even if it fluctuates and you have some periods where there's a smaller impact on your daily life, you might have this impairment for your lifetime.

Find out more about [impairment categories](#), and what they mean.

It doesn't matter what caused your impairment. For example, if you've had it from birth or acquired it from an injury, accident or health condition.

It also doesn't matter if you have one impairment, or more than one impairment.

Is your impairment permanent or likely to be permanent?

To meet the disability requirements, we need to know that your impairment is or is likely to be permanent.²¹ Permanent under the law for the NDIS means enduring. This means we need to know whether your impairments are expected to be lifelong and not likely to go away or significantly improve over time.

We'll focus on your impairments and not on the cause of your impairments, or your disability or diagnosis.

You might have some times in your life where there's a smaller impact on your daily life, because your impairment may be episodic or fluctuate in intensity.²² Your impairment can still be permanent due to the overall impact on your life, and the likelihood that you'll be impacted across your lifetime.

Even when your condition or diagnosis is permanent, we'll check if your resulting impairment is permanent too. For example, you may not be eligible if your impairment is temporary, or if there are known, available and appropriate evidence-based clinical, medical or other remaining treatments options that are likely to remedy the impairment.

Generally, we'll consider whether your impairment is or is likely to be permanent after all known, available and appropriate treatment options have been pursued.

If you give us evidence you have been diagnosed with a condition on [List B](#), we'll likely decide your disability is from an impairment that's likely to be permanent.

Is there any medical treatment for your impairment?

We don't fund supports to treat your impairment. Supports including the diagnosis, early intervention and treatment of health conditions, including ongoing or chronic health conditions, are funded by the Australian health system instead of the NDIS.

Your impairment will likely be permanent if your treating professional tells us there are no further treatments that could remedy it.

Your treating professional will tell us or be asked to certify if there are medical, clinical or other treatments that are likely to remedy your impairment. We need to understand whether there are treatments that are:²³

- **known** – the treatment can be identified by an Australian medical practitioner as a suitable treatment for your impairment.
- **available to you** – we need to take account of whether there are genuine barriers that prevent you from accessing treatment including, but not limited to, the nature of your impairment and your ability to access treatment.
- **appropriate for you and your impairment** – we need to consider whether the treatment could remedy your impairment and is suitable and safe for you to undergo. Your ability to undergo treatment will be assessed according to your capabilities, your health and other personal circumstances, including your living arrangements.
- **evidence-based** – there's proof the treatment is likely to be effective.

When we look at what treatments are available to you, we think about whether the treatment is suitable for your personal situation. The word treatment should be understood in the broadest sense and may include changes to your diet and lifestyle.

If you're still undergoing or have recently had treatment, we may not be sure you have a permanent impairment if that treatment could remedy the impairment.²⁴

In some situations, it may be clear your impairment is likely to be permanent while you're still undergoing treatment or rehabilitation. For example, you may still need treatment and rehabilitation for a spinal cord injury, but it's clear you're likely to have a permanent impairment.

You might still have a permanent impairment, even if its effects may change over time.²⁵

For degenerative impairments, or those that get worse over time, we consider them permanent if treatment isn't likely to remedy the impairment. That is, the treatment won't cure the impairment or come close to removing its effects.

Does your impairment substantially reduce your functional capacity?

The laws for the NDIS describe **functional capacity** in relation to an activity, as a person's ability to do the activity without **help from others**, **assistive technology** or **modifications**, or only with assistance from commonly used assistive technology or modifications.

Commonly used assistive technology or modifications are the things you would usually use to do an activity. For example, glasses, walking sticks and hearing aids.²⁶

When we think about help or assistance a child needs to do an activity, we look at whether the help the child needs from others is similar to other children of the same age.²⁷

When we consider a person's functional capacity we exclude, as much as possible, the impact of a person's environmental and personal circumstances.²⁸

Your permanent impairment needs **to substantially reduce your functional capacity** in relation to one or more of the following activities, considering each activity as a whole:

- **Communicating** – how you speak, write, or use sign language and gestures, to express yourself compared with other people your age. We also look at how well you understand people, and how others understand you.
- **Socialising** – how you make and keep friends, or interact with the community, or how a young child plays with other children. We also look at your behaviour, and how you cope with feelings and emotions in social situations.
- **Learning** – how you learn, understand and remember new things, and practise and use new skills.
- **Mobility, or moving around** – how easily you move around your home and community, and how you get in and out of bed or a chair. We consider how you get out and about and use your arms or legs.
- **Self-care** – personal care, hygiene, grooming, eating and drinking, and health. We consider how you get dressed, shower or bathe, eat or go to the toilet.
- **Self-management** – how you organise your life. We consider how you plan, make decisions, and look after yourself. This might include day-to-day tasks at home, how you solve problems, or manage your money. We consider your mental or cognitive ability to manage your life, not your physical ability to do these tasks.

Your impairment substantially reduces your functional capacity if you usually need disability-related supports to participate in or complete the above tasks.²⁹

These disability-related supports include:

- a high level of support from other people, such as physical assistance, guidance, supervision or prompting
- assistive technology, equipment or home modifications that are prescribed by your doctor, allied health professional or other medical professional.

To help us decide if you're eligible, we need to know your capacity and where you need more help. We get this information from you when you apply to the NDIS.

If you have more than one permanent impairment, we'll consider them together, to see if they substantially reduce your functional capacity.

Your needs might go up and down each day or each month. We consider your ability over time, taking into account your ups and downs. For example, someone with relapsing remitting Multiple Sclerosis (MS) may have varying levels of support needs across a day, week or month.

How does a child's impairment affect their daily life?

To help us decide if a child's ability is substantially reduced, we compare their abilities with other children of the same age.

If a child's ability is much less than most other children the same age, they may meet the disability requirements. For example, if they:

- need assistive technology, equipment or home modifications to participate in daily activities – except for common items like glasses
- usually need more help to join activities.

We'll look at the early intervention requirements if:

- a child's impairment doesn't substantially reduce their ability right now, but might in the future
- a child's impairment currently substantially reduces their ability, but may not after receiving supports.

Learn more about the [early intervention requirements](#).

What if you have a hearing impairment?

Some hearing impairments may lead to a substantially reduced functional capacity.

We look at the information and evidence you've given us to decide if your hearing impairment leads to a substantially reduced functional capacity. However, we'll generally decide you have a substantially reduced functional capacity if your hearing loss is at least 65 decibels in your better ear. This is based on a pure tone average of 500Hz, 1000Hz, 2000Hz and 4000Hz.

We may also decide you have a substantially reduced functional capacity if your hearing loss is less than 65 decibels in your better ear. We may decide this if either:

- you also have another permanent impairment, such as a vision or cognitive impairment
- you give us evidence your speech detection and speech discrimination outcomes are significantly poorer than expected.

Does your impairment affect your social, work or study life?

We look at how your impairment affects your ability to work, study or take part in social life.³⁰ This means your permanent impairment affects how you can find and keep a job, contribute to your community, or join social activities. We get this information from you when you apply to the NDIS.

We look at your ability to do things like:

- find and keep a job, or start your own business
- study
- play sport
- go to the movies
- volunteer
- participate in cultural activities or ceremonies
- travel.

You can meet this criteria when your ability to work, study or socialise is affected by your impairment in some way.

Will you likely need NDIS supports for your lifetime?

You must be likely to need NDIS supports for your lifetime.³¹ [NDIS supports](#) are the services, items and equipment that can be funded by the NDIS.

NDIS supports are investments that help you build or maintain your functional capacity and independence, and help you work, study or take part in social life.

Even if your needs go up and down over time, or happen episodically³², we may still consider it's likely you'll need NDIS supports for your lifetime.³³

We consider your overall situation to answer this question.

When we decide if you'll likely need NDIS supports for your lifetime, we consider:

- your life circumstances
- the nature of your long-term support needs
- whether your needs could be best met by the NDIS, or by other government and community services.

For example, you may have an impairment that is caused by a chronic health condition. Many chronic health conditions are most effectively managed or remedied through medical management through the health system. If this is the case, we may decide that you don't need NDIS supports for your lifetime.

Learn more about [reasonable and necessary supports](#) and [NDIS supports](#).

Do you need early intervention?

Early intervention is usually early access to support, to reduce your future needs for disability-related supports.

Early intervention can be for both children or adults and may only be needed for a short time. You won't need these supports for your lifetime, so your treating professional or your early childhood partner will tell us how early intervention support could benefit you or your child.

You'll meet the early intervention requirements if you meet all the following:

- You have an [impairment that is or is likely to be permanent](#).
- [Early intervention supports will likely benefit you](#), and it means you'll need less disability support in the future.
- [Early intervention supports benefit you by](#) reducing the impact of your impairment on your functional capacity in relation to one or more activities. This means your ability to communicate, socialise, learn, move around or look after yourself and organise your life. Or the supports prevent your functional capacity from getting worse, improve your functional capacity or support your informal supports.
- [The early intervention supports you need are NDIS supports](#).

There are different requirements for [children younger than 6 with developmental delay](#) to meet the early intervention requirements.

If we have evidence a child younger than 7 has been diagnosed with a condition on [List D](#), we'll decide they meet the early intervention requirements.

You may also meet the early intervention requirements if you're [aged between 0 and 25 with a hearing impairment](#).

We also need to understand how NDIS supports benefit you, like building your skills and increasing your capacity, so that you may no longer need NDIS supports. If you meet the early intervention requirements, your support needs are more likely to change, and you may only need NDIS supports for a short time. We'll regularly check your eligibility when we reassess your plan, and at other times too.

If you no longer meet the early intervention requirements, we'll check if you meet the disability requirements. Learn more about how we check if you're [still eligible for the NDIS](#).

Is your impairment permanent or likely to be permanent?

To meet the early intervention requirements, there must be enough evidence that you have at least one of the impairments below and your impairment is or is likely to be permanent.³⁴

An impairment is a loss of or damage to the body's structure and/or how the body functions.

We will look at:

- your body's functions
- your body structure
- how you think and learn.

An [impairment](#) could be:³⁵

- **intellectual**
- **cognitive**
- **neurological**
- **sensory**
- **physical.**

You may also be eligible for the NDIS if you have a **psychosocial disability**.³⁶ This means you have reduced capacity to do daily activities and tasks due to your mental health. Your psychosocial disability might vary at different times in how much it impacts your daily life. Even if it fluctuates and you have some periods where there's a smaller impact on your daily life, you might have this impairment for your lifetime.

Find out more about [impairment categories](#), and what they mean.

When we decide if your impairment is likely to be permanent, we consider the same things as in the [disability requirements](#).

We'll need evidence that at least one of your impairments will be permanent, or likely to be permanent.³⁷

If you give us evidence you have been diagnosed with a condition on [List B](#), we'll decide you have an impairment that's likely to be permanent.

For some very young children, problems in body function can't be easily measured. However, the child may be at risk of a future diagnosis and permanent impairment. A health or allied health professional will need to provide evidence about the child and if they are likely to have a permanent impairment.

Will early intervention benefit you?

We need to decide that getting early intervention supports means you'll likely need less disability-related supports in the future.³⁸

How will early intervention benefit you?

We need to know that early intervention supports will help you with at least one of the following:³⁹

- addressing the impact of your impairment on your ability to communicate, socialise, learn, move around, look after yourself and organise your life
- preventing your functional capacity from getting worse
- improving your functional capacity
- supporting your informal supports, which includes building their skills to help you.

To help us decide if the early intervention will help you in these ways, we look at all of the following:⁴⁰

- how your impairment might change over time
- how long you've had your impairment
- if there's been a significant change to your impairment
- what supports you have received
- any supporting evidence you give us.

Will the early intervention supports you need be NDIS supports?

The early intervention support that you would likely benefit from must be NDIS supports.⁴¹

Learn more about [reasonable and necessary supports](#) and [NDIS supports](#).

What about people aged between 0 and 25 with a hearing impairment?

If you're aged between 0 and 25 with a hearing impairment, you may meet the early intervention requirements. We'll decide this if you give us evidence of all of the following:

- You're aged between 0 and 25.
- You have auditory neuropathy or hearing loss of at least 25 decibels in either ear at 2 or more adjacent frequencies – see below.

We need evidence of your auditory neuropathy or hearing loss from a specialist audiological assessment. The assessment might include electrophysiological testing when required. The evidence must show your hearing loss is or is likely to be permanent.

If you're aged 26 or older with hearing loss, we'll check if you're eligible in the same way we consider all other impairments. You may be eligible under the [disability requirements](#).

What about children younger than 6 with developmental delay?

Children younger than 6 with developmental delay may be eligible for the NDIS under the early intervention requirements.⁴²

Developmental delay is a term used to describe a delay in a child's development. It means that a child finds it much harder to do everyday things that other children their age can do, for example, dress themselves, talk or walk. A child with developmental delay needs lots of extra help to do everyday things compared with other children their age.

We'll need to know the child:

- is younger than 6 on the day we decide whether they're eligible⁴³
- has [developmental delay](#)⁴⁴
- will benefit from early intervention supports that are NDIS supports.

An early childhood partner can also provide supports to children who aren't eligible for the NDIS.

Learn more about the [early childhood approach](#) and [early connections](#).

Does the child have developmental delay?

When we decide if a child has developmental delay, we use the definition in the law for the NDIS.⁴⁵

We need to know the delay:

- is [due to mental or physical impairments](#) or a combination of both
- [substantially reduces the child's functional capacity](#) in relation to one or more major life activities compared with other children their age
- means [the child needs specialist services](#) from more than one professional working as a team to support the child for an extended time. Usually for longer than 12 months.

Is the delay due to mental or physical impairments?

First, we need to know the developmental delay is due to a mental or physical impairment, or a combination of mental or physical impairments.⁴⁶

An impairment is a loss or significant change in at least one of:

- the child's body functions
- the child's body structure

- how the child thinks and learns.

Parents, carers, early childhood partners and other professionals can understand the child's body function by:

- observing their activities during play and daily tasks
- comparing their activities to other children of the same age.

Does the delay substantially reduce the child's functional capacity?

We need to know the delay substantially reduces the child's functional capacity compared with other children their age.⁴⁷

This means the child has a significantly lower ability to do everyday activities, when compared with other children the same age. Or the child does things in a significantly different way to other children their age because of their reduced capacity.

The child would also need much more support to do the activity, compared with other children their age.

The substantial reduction in functional capacity must be in relation to at least one of the following areas of major life activity:

- **Self-care** – how children take care of themselves, shower, bathe, dress, eat, drink, toilet, groom, and sleep.
- **Receptive and expressive language** – this involves skills such as gesture, sign language, listening, giving and receiving information, communicating wants and needs through facial expressions, vocalisations or speech, and interaction with others.

A substantial reduction in functional capacity for either receptive language or expressive language will meet the criteria – it doesn't need to be both.

- **Cognitive development** – learning and applying knowledge. This includes areas such as:
 - understanding and remembering information
 - attention
 - learning new things
 - practising and using new skills
 - planning and making decisions
 - problem solving
 - developing pretend play skills
 - developing play interests

- emotional and sensory regulation
 - developing emotional intelligence
 - social awareness
 - safety awareness.
- **Motor development** – this includes participation in everyday activities like moving around the home and community and manipulating objects.

We need evidence from a health, allied health or early childhood professional, who uses multiple sources of information about the child's ability to do everyday activities.

This will include information that parents or carers report about their child. It will also include a mix of standardised assessments of developmental and functional capacity, both in everyday activities and settings.

It should also include observations in everyday play, learning, activities or routines to better understand how the child participates in these everyday activities.

For very young children where functional capacity can't be measured, the child may be eligible if there's significant risk of a future diagnosis or developmental delay. We need evidence of this from a health or allied health professional's informed clinical opinion.

Does the child need specialist services from more than one type of professional for an extended time?

We need to know that the child needs a mix of specialist care, treatment or other services, due to their developmental delay. The child must also need these services for an extended duration. For example, longer than 12 months.⁴⁸

We need evidence from an early childhood professional, such as an early childhood teacher, educator or allied health professional who knows the child. The evidence must show the child needs all of the following:

- **A service response that involves more than one professional working as a team to support the child.** This means the child needs support for multiple activities, and across multiple everyday settings such as the home, community and early childhood centres. The child must need more support than what's expected for a child the same age.
- **A team that works collaboratively, by communicating and sharing information, knowledge and skills.** The support must be individually planned and coordinated. The team will build the capacity of the child's family and other important people in the child's life, such as carers, educators and professionals, about the child's individual

needs. This support should be embedded in everyday play, learning, activities and routines.

- **More support than an individual discipline providing a unilateral response to a single problem.** This means the child needs support from more than one professional supporting one area of delay. This is known as interdisciplinary care. For example, a child is unlikely to be eligible if a speech pathologist alone can help their language delay, without needing support or consultation from other professionals.
- **Supports for an extended duration.** This means a health, allied health or early childhood professional who knows the child determines they need support for an extended time. Usually longer than 12 months. A child will likely meet this criteria if there's clear evidence that they'll need early intervention support for more than 12 months.

Some children in remote areas might not have access to a team of professionals. If so, they may still be eligible if the one professional needs to provide the supports to the child across multiple activities and across multiple everyday settings.

The professional should consider multiple sources of information, including:

- parent or carer reports
- a mix of standardised and culturally appropriate developmental or functional assessments in everyday activities and everyday settings
- observations in everyday play, learning, activities, and routines.

How do we work out if the child meets the criteria for developmental delay?

We'll need a range of information about the child, observed in everyday activities and settings they usually participate in. This should include parent or carer reports and standardised assessments of developmental and/or functional capacity.

Early childhood partners are early childhood professionals who give us evidence of developmental delay to help us decide if the child is eligible. An early childhood partner will meet with children and families to better understand the child's day to day life, and any concerns about their development.

Early childhood partners will observe a child in everyday settings like home and childcare and may complete assessments using screening tools. This information helps us decide if a child meets the early intervention requirements for developmental delay.

Parents and carers can also provide copies of existing reports, assessments or letters about the developmental delay, if they have them.

We may also ask for evidence from a variety of sources, including mainstream services. For example, we may also ask for evidence from your doctor, child health nurse, or other most appropriate treating professional.

What if there are no early childhood partners in your area?

If there are no early childhood partners in the child's area, a mainstream, community, or health service, for example an Aboriginal medical service or community controlled organisation, can give us a report for evidence of developmental delay.

Will the child's early intervention supports be NDIS supports?

To meet the early intervention requirements, the supports must be NDIS supports.⁴⁹. Find out more about [reasonable and necessary supports](#) and [NDIS supports](#).

What happens if a child with developmental delay is eligible?

If we decide a child with developmental delay is eligible for the NDIS, they'll become a participant. But they're usually no longer eligible after they turn 6.

This is because they'll no longer meet the early intervention requirements under developmental delay. To remain an NDIS participant after they turn 6, the child will need to have an impairment that is or is likely to be permanent and meet the requirements for [disability](#) or [early intervention](#).

We'll talk to parents or carers before a child turns 6 and explain what information we need to decide if the child is still eligible. Learn more about [leaving the NDIS](#).

Example

Hunter is 5 years old. When he was 3, Hunter became an NDIS participant under the early intervention requirements for developmental delay.

While Hunter is a participant we talk to his parents about his progress and outcomes from the early intervention supports received. We listen to understand how he has built his capacity to work towards his goals. Hunter's family have also built their capacity to support his development. They're happy with the outcomes achieved. He is building his communication and social skills and as a result is needing less early intervention support. We also help Hunter and his family connect to other government and community services.

Hunter's parents understand that he may no longer be eligible for the NDIS after he turns 6. We explain the disability and early intervention requirements, including that if Hunter is to continue to be eligible for the NDIS after he turns 6, we need evidence that he has an impairment that's likely to be permanent. Before Hunter turns 6, we let Hunter's parents know what information is needed to decide if he would continue to be eligible under the [disability](#) or [early intervention](#) requirements after he turns 6.

Learn more about [leaving the NDIS](#) and [mainstream and community supports](#).

What if a child doesn't meet our criteria for developmental delay?

Early childhood partners offer supports to children younger than 6 who don't meet the early intervention requirements for developmental delay.

A child may have **developmental concerns**. This means a child younger than 6 is developing slower compared with other children their age, but the delay doesn't meet our definition for developmental delay.

For example, a child's functional capacity may be substantially reduced in one or more areas. But it's unclear if the child needs support from a team of professionals for an extended duration.

An early childhood partner can offer **early supports** to children younger than 6 with developmental concerns. They can also help the child and family connect to other government and community supports.

Learn more about [early connections](#).

How do you apply to the NDIS?

Applying to the NDIS is how you let us know you want to become an NDIS participant.

If you're aged 9 and older, there are a few ways you can apply:⁵⁰

- Your local area coordinator can help you apply. They can help you through the application process and be your point of contact. [Find your nearest location](#).
- Sometimes you may not have a local area coordinator in your area. You can contact us on 1800 800 110 to discuss other options available to you.
- First Nations people can call our First Nations Connect service on 1800 411 640, for a culturally respectful and tailored experience.

For children younger than 9, we encourage parents and carers to talk to an [early childhood partner](#) first about their child's development. They can offer supports to children before they apply and let parents and carers know if the NDIS is right for their child.

When you apply, you or your authorised representative will need to:

- give us the information and any documents we need to confirm your identity. Learn more about [evidence of identity](#) and [privacy](#)
- give us the information and any documents we need to decide if you're eligible⁵¹
- sign or certify the NDIS application⁵²

- talk about your needs and current situation.

When we talk to you, we'll listen to understand what is important to you. We'll also ask questions to make sure we know all the ways we can help.

We can use this information to help you connect to other supports if you want us to. Learn more about [community connections](#) and [early connections](#).

Other people can help you apply if you want them to. Sometimes they can apply on your behalf. Learn more about [who can help you apply](#) and about [how to apply to the NDIS](#) in the [Applying to the NDIS factsheet](#).

What information do we need in your application?

Evidence of your identity

The [Evidence of identity factsheet](#) shows what information we need to confirm your identity. When you apply for the NDIS, you'll need to give us copies of these documents. If you can't do this, let us know so we can work out what to do depending on your situation. We'll still need to confirm your identity before progressing your application.

To show us you're younger than 65 when you apply, live in Australia, and that you're an Australian citizen, permanent resident or hold a Special Category visa that is a protected special category visa, you can give us either of the following:

- consent to access and use your Centrelink record
- copies of documents or other evidence that we ask for if you apply in person or over the phone.

In most cases, we can just use your identity documents.

If you don't have the identity documents we need, let us know. We'll talk with you about your situation and work out the best way to support you.

Evidence of your impairments?

We need evidence of your impairments, to help us work out if you're eligible. To get information about this, ask your [appropriate treating professional](#). For children younger than 6 with developmental delay, an [early childhood partner](#) can give evidence of developmental delay. Your treating professional or early childhood partner can contact us if they need to discuss what evidence to provide.

Your treating professional might be your doctor, specialist, or allied health service provider. You should use a professional who:

- has worked with you for a long time, usually for at least 6 months

- is the [most appropriate type of professional](#) to give evidence about your impairment
- is qualified and registered in their area of practice with the [Australian Health Practitioner Regulation Agency](#) or relevant professional authority.

If your treating professional doesn't meet these requirements, we may not be able to confirm the information in your application. If we need more evidence we may need you to give us further information.⁵³

When we check if you're eligible for the NDIS, we mainly think about the information you give us when you apply.

Learn more about who can give us evidence on the following pages:

- [How to gather evidence](#)
- [Guide to the NDIS for GPs and health professionals](#).

You can also learn more about [how we use, collect and store your personal information](#).

What if you're in a remote or very remote area?

If you live in a remote or very remote area we understand it can be hard to get the required evidence from an [appropriate treating professional](#). If it's hard to get your treating professional to do this, let us know.

We use a technical definition for remote and very remote. You'll need to live in an area that's classified as MM6 or MM7 on the [Modified Monash Model](#) to be considered remote or very remote. The Modified Monash Model is a mapping tool we use to figure out how remote an area is. MM6 is the category for remote communities, and MM7 is the category for very remote communities.

If you live in a remote or very remote area where you can't access an appropriately qualified treating professional, an Aboriginal medical service or Community controlled organisation can help you gather the information we need. Sometimes, we'll reach out to you to help you get the required evidence and support you to apply.

Who can help you apply?

You can ask someone to help you apply if you want to. They can help you:

- make your decision to apply to the NDIS
- gather the information we need.

You can choose who helps you. For example, you could ask for help from:

- a family member
- a friend

- a carer
- a partner
- a support worker or service provider
- staff in a residential aged care facility
- an Aboriginal medical service
- a community controlled organisation
- your treating professional
- hospital staff.

With your permission, we can share information with these people during your application. For example, they could call us to check how your application is progressing. You can let us know if you would like us to share information. Learn more about [consent](#) and [how to give consent](#).

Can someone else apply for you?

Sometimes, another person may have legal authority to apply to the NDIS on your behalf.

If you're younger than 18, the person or people with parental responsibility for you will apply for you.⁵⁴ This is often your parents or legal guardian. In some situations, we can decide someone else is your representative for NDIS matters.⁵⁵ Learn more about [child representatives](#).

If you're an adult, these people may be able to apply to the NDIS on your behalf:

- a person you give consent to act as your authorised representative – this means you give them permission to apply for you
- your guardian
- a person with power of attorney who can make personal and health decisions for you
- a person with advance care health directive.

If you're an adult and want someone else to apply for you, you can tell us in person, or over the phone. We'll need to make sure you agree to someone else applying for you.

If you have a guardian or a power of attorney, we'll ask for information that shows the person is your guardian or power of attorney.

How do we check your application?

Before we can accept your application, we make sure it's been made by the right person. That is, the application is from you, or [someone who can apply for you](#).

We then check all the information we need has been provided.

What if you haven't given us enough information with your application?

If you don't have all the right information, we'll help you work out what to do. You can also let us know if there's a mistake. We can work with you to help complete the application properly. We can't decide if you're eligible until we have a complete application.

Once you have completed your application with all the right information, we'll check whether you're eligible. That is, we'll check that:

- you meet the age and residence requirements
- you meet the requirements for disability, early intervention, or both.

[If we still need more information](#) to check you're eligible, we'll let you know what you need to give us.

Learn more in the [Applying to the NDIS factsheet](#).

What if we need to verify the evidence in your application?

When we check your application, we may at times ask you for more information.

If we believe you or someone else has relevant information we need, we'll make a request in writing for more information.

The request will explain:

- what information we need
- how and when to provide it
- your legal rights and responsibilities. This includes your right to not provide information if doing so may result in a penalty.

We understand people can make mistakes, so we want to work with you to get the right information to help us make a decision.

How do we decide if you're eligible?

Once we have your completed application, we review all the information we have in your application.

This will help us decide if you're eligible for the NDIS. As part of the process, we'll also need to check your identity.

You're eligible for the NDIS if you meet the requirements for:

- age

- residence
- [disability](#) or [early intervention](#) or both.

If you don't meet either the disability requirements or the early intervention requirements, you won't be eligible for the NDIS. An [early childhood partner](#) or [local area coordinator](#) can help you connect with other government and community supports.

When will we decide if you're eligible?

Once we have your complete application, we'll decide one of the following within **90 days**:

- [you're eligible for the NDIS](#)
- [you're not eligible for the NDIS](#)
- [we need more information](#).

The amount of time we have to make one of these decisions depends on the date we determine your application is complete. We'll send you a letter telling you the date we need to make a decision by. This won't be longer than 90 days.⁵⁶

If we've sent you a letter asking you for more information, we have **14 days** once you give us the information we need, to make a decision about your application or request more information again.

Learn more about what you can expect from us and what we consider when we receive your application in our [Participant Service Charter](#).

We can make a decision quicker in urgent circumstances. Let us know if your situation is urgent, for example, if you're about to leave a hospital or custodial setting. Learn more about our [timeframes for urgent decisions](#).

How do we consider your evidence of disability or developmental delay?

When we're deciding if you're eligible, we may look at things like:

- how old your evidence is
- who provided your evidence.

If we get more than one type of evidence from you, we might consider some evidence over others. We call this [weighing evidence](#).

What if we need more information to decide if you're eligible?

When we decide if you're eligible, we look at:

- the information in your NDIS application

- any other information we have.

We need enough information in your application to show us you're eligible for the NDIS by meeting the requirements for disability or early intervention.

Sometimes we might need to ask you for more information. For example, we may not have enough information about your functional capacity.

We'll ask you for more information if we need it to make sure we have the full picture.⁵⁷

We might ask you for more information if:

- your application doesn't have all the information we need
- we need to answer a particular question.

We'll only ask for more information if we need it to decide if you're eligible.⁵⁸ If we need more information, we'll let you know:

- what you need to do
- what information we need
- when you need to give us the information.

If we ask for more information, you'll have at least **90 days** to give it to us. We can't decide if you're eligible until we have this.

If we can't contact you within **90 days**, or you don't give us the information within the timeframe, we take this to mean you've withdrawn your application and we'll stop processing your application.⁵⁹

You can ask for more time if you need it. We can give you more time if we think it's reasonable for your situation.

If you don't get the information to us in time, you can apply again.

What happens after we get your information?

Once you give us the information we need, we then have **14 days** to decide if:⁶⁰

- you're eligible for the NDIS
- you're not eligible for the NDIS
- we need more information – for example, if the information you gave us isn't what we need.

Learn more about what you can expect from us and what we consider when we ask for more information in our [Participant Service Charter](#).

What happens if we don't decide on time?

If we don't meet our decision-making timeframes, we have to treat this as if we decided you're not eligible.⁶¹

If this happens, we'll automatically review the decision that you're not eligible.⁶² You don't need to do anything.

We'll then review your application again. We'll contact you to let you know the outcome.

Learn more about [reviewing our decisions](#).

What happens after we decide?

What happens if you're eligible?

On the day we decide you're eligible for the NDIS, you become an NDIS participant.⁶³

The time that you remain eligible for the NDIS depends on the eligibility requirements you met, your individual circumstances and support needs.

You'll remain an NDIS participant as long as you continue to meet the eligibility requirements for the NDIS. This means you'll need to continue to live in Australia and be an Australian citizen, permanent resident or hold a Special Category visa that is a protected special category visa. You'll also need to continue to meet requirements for [disability](#) or [early intervention](#). Learn more about [whether you'll always be eligible](#) and [leaving the NDIS](#).

We'll send you a letter to let you know:

- you're eligible
- if you met the requirements for disability, early intervention, or both⁶⁴
- the next steps.

Your letter will also confirm the date you became eligible for the NDIS.⁶⁵

Your [impairment categories information](#) will be listed in the letter you receive which tells you your application to the NDIS has been successful.⁶⁶

What do we mean by impairment categories information?

This information tells you the impairment categories that cover the impairments for which you meet the requirements for disability, early intervention or both. Under the NDIS law, this is called a notice of impairment. We use the word impairment because the laws for the NDIS are focused on impairments, and not on a disability or diagnosis.

Participants who have the same diagnosis may have different impairments and have different support needs. This means that participants with the same diagnosis may need different NDIS supports.

Impairment categories are the way we group together different types of impairments you have. This helps us understand how your disability impacts your everyday life.

Your **impairment categories information** tells you about the impairment category or categories that cover the impairment(s) that meet the requirements for disability, early intervention or both. It doesn't list your specific impairments or conditions.

We look at your evidence of disability to decide the impairment categories that cover your impairments that meet the disability and/or early intervention requirements. We include your impairment categories information in the letter which tells you your application to the NDIS has been successful.

The impairment categories don't apply to children younger than 6 who are eligible under the early intervention requirement for developmental delay only. Instead, they'll have **developmental delay** listed in their letter.

What are the impairment categories?

Below is a detailed explanation of the **6** impairment categories. It includes some examples of related conditions, but it's not a full list of conditions. Depending on your circumstances, one or more categories may apply.

These category descriptions were developed through a review of Australian and international research, as well as through consultation with Australian disability peak bodies.

The categories are:

- **intellectual** – differences in intellectual function that impact how you think, plan and learn, or how you make judgements, problem solve and reason. These differences may impact your everyday life skills. Intellectual impairment is usually identified in childhood and continues throughout life. While intellectual impairment can be genetic, sometimes the cause is unknown. Conditions such as Down syndrome and severe intellectual disability are examples of conditions known to affect intellectual function.
- **cognitive** – differences in body functions related to how you think, understand or remember. These differences may impact your alertness, attention, orientation, information processing, communication, learning, memory or problem solving. Cognitive impairments can be present at any age. Conditions such as brain tumour and traumatic brain injury are examples of conditions known to affect cognitive function.

- **neurological** – differences in body functions related to the nervous system. Your nervous system includes your brain, spinal cord and nerves. Changes to these parts of the body can impact movement, sensation and speech, as well as processing of sensory information. Conditions like Parkinson's disease, multiple sclerosis and Autism are examples of conditions known to affect neurological capacity. Autism is included in this category as it's a condition that affects the way your brain processes information. Autism usually becomes noticeable during childhood development.
- **sensory** – differences in the structure and or functions of the eyes and ears. These differences may impact your sight and hearing. Differences to the structure of your eyes and ears might be present at birth (congenital) or acquired later in life. Conditions like vision loss or vision impairment, hearing loss or deafness; or deafblindness are examples of conditions known to affect sensory functions.
- **physical** – differences in your body's functions that impact on your ability to move parts of your body. These differences may impact body movements, balance or coordination. They may impact on doing daily activities like walking, going from sitting to standing, getting dressed, showering, speaking or eating and drinking. Your physical differences may affect how quickly your body gets tired. Conditions such as amputation of a limb or limb differences, and rheumatoid arthritis are examples of conditions known to affect physical functions.
- **impairments relating to a psychosocial disability** – differences in the way you complete daily activities because of your mental health. These differences may impact your ability to think clearly, manage everyday tasks, get an education, find and keep a job, live independently or maintain relationships. Conditions like Bipolar affective disorder or Schizophrenia are examples of conditions known to affect psychosocial function.

Example

Oscar is 25 years old and is eligible for the NDIS under the age, residence and disability requirements. Oscar has a permanent impairment relating to his spinal cord injury.

Oscar gets a letter telling him his NDIS application was successful, and he's now a participant. This letter includes his impairment categories information.

The letter tells him the impairment categories that relate to his permanent impairments are physical and neurological.

At Oscar's first planning conversation, they discuss the supports that relate to Oscar's physical and neurological impairments. Oscar understands that he can ask to change his impairment categories information if needed.

How do we decide your impairment categories

26 August 2026

Applying to the NDIS

Page 30 of 57

This document is correct at the date of publication.

Always visit ourguidelines.ndis.gov.au for the latest version.

To work out your impairment categories, we look at the information and evidence you gave us when you applied for the NDIS. We make sure the impairment categories cover your impairments that meet the requirements for disability, early intervention or both.

You may have multiple impairments for which you meet the disability, and/or early intervention requirements. We consider all permanent impairments that result in you having substantially reduced functional capacity, and/or the need for [early intervention](#). Your impairment categories information may include one or multiple impairment categories.

More impairment categories doesn't always mean you need more NDIS support. If you have multiple impairments which relate to a single impairment category, then we'll record one impairment category.

The same impairment can affect people in different ways. When we decide your impairment categories, we think about how your impairments affect your life and the way your body functions.

Example – multiple impairments covered by the same impairment category

Mei was diagnosed with fusion of the cervical spine after birth. Later in life, Mei had a below-knee amputation.

Mei gives us evidence showing permanent damage to their body functions. This is due to limited range of motion in the neck and upper spine, and the partial loss of a lower limb.

Although Mei has two separate impairments, Mei's impairment categories information will include only the physical impairment category. This is because both the cervical spine and below-knee amputation are types of physical impairments.

Can I ask to change my impairment categories information?

You can ask to change your impairment categories information⁶⁷ if you think the impairment categories in your letter are not correct.

[Let us know](#) if you want to change your impairment categories. You'll need to give us information about what impairment categories may need to change and why. Please give us evidence of any new impairments you have.

We'll look at the evidence against the disability or early intervention requirements to work out if we need to change your impairment categories. We'll also consider the functional impact of your impairment alongside the information and evidence we have.

We'll let you know if we decide to change your impairment categories or not. If we do change them, you'll get a letter with new impairment categories information.⁶⁸

If we decide to change, or not to change, your impairment categories information and you don't agree with our decision, you can ask for a review of the decision. Learn more about [reviewing our decisions](#).

Example – decision to add an impairment category

Omar meets the disability requirements for an intellectual disability.

Omar's impairment categories information states intellectual and cognitive to cover the impairments they meet the disability requirements for, associated with their intellectual disability.

Omar's nominee, Shae contacts the NDIA to advise that Omar has recently been diagnosed with Parkinson's disease. Shae asks to change Omar's impairment categories information to reflect this.

As part of the request, Shae provides diagnostic information from Omar's neurologist. This evidence demonstrates Omar's Parkinson's disease results in a permanent impairment that impacts his neurological function and has a substantial impact on his functional capacity.

We consider the application and evidence and decide to change the impairment categories information. This is because Omar's Parkinson's disease results in a permanent impairment which causes substantial reduction in functional capacity.

Omar is sent a letter to explain we have changed his impairment categories information. It now also includes the neurological impairment category, along with the intellectual and cognitive impairment categories.

Example – decision to not change impairment categories information

Farrah has a vision impairment. Her impairment categories information includes one category – sensory. The sensory category covers her vision impairment.

Farrah contacted us to let us know she broke her arm. She has been receiving treatment for her broken arm from the health system, but it's taking a long time to heal. Farrah applies to change her impairment categories. She would like to add the physical category. This is because while her arm is still healing, she has reduced range of movement in her arm and difficulties carrying out some everyday tasks.

We look at Farrah's application and decide not to add a new category. Her current category of sensory is still correct. We make this decision because there's no evidence that the impairment caused by Farrah's broken arm is permanent or likely to be permanent. This means it doesn't meet the requirements for disability or early intervention.

Farrah gets a letter to tell her about the decision. Farrah knows that she can request an internal review of this decision if she wishes.

How will we create your first plan?

After you receive the letter confirming you're eligible, we'll start making your plan within **21 days**. Find out more about [creating your plan](#).

If you received help from a partner to apply to the NDIS, we can build on the information and goals we talked about and include these supports in your plan.

You'll receive a plan that sets out your NDIS supports. [NDIS supports](#) are the services, items and equipment that can be funded by the NDIS. The supports must relate to your disability.⁶⁹ This means there must be a direct link between your disability support needs and the NDIS supports we fund.

Our [Participant Service Guarantee](#) says we must approve your first plan within **56 days** after you become a participant.

For more information, check out [creating your plan](#) and [reasonable and necessary supports](#).

Will you always be eligible for the NDIS?

There are many reasons for leaving the NDIS.

Some people decide they don't want to be a participant anymore.

You'll also leave the NDIS if you're no longer eligible.

When we [reassess your plan](#), we check that all your details are correct and up to date. We also look at any new information we have received.

If you're eligible under the **early intervention requirements**, your support needs are more likely to change. We'll check at each plan reassessment and at other times, whether you still meet the early intervention requirements.

For example, during a plan reassessment, evidence may show you no longer meet the early intervention requirements because you have built your skills and capacity and will no longer benefit from NDIS supports.

If you're eligible under the disability requirements, we don't expect your disability to change and you'll likely need NDIS supports for your lifetime. We'll only ask you for more information about your eligibility if there's evidence that you may no longer meet the disability or residence requirements.

You can find out more about the eligibility requirements, and how we check these, at [How to check your eligibility](#). [Children with developmental delay](#) will usually leave the NDIS after they turn 6.

Over time, you might develop your skills and independence and not need NDIS supports anymore.

If you met the requirements for early intervention and not disability, you usually won't be eligible after the early intervention supports have benefitted you. For example, if you needed NDIS supports to achieve your goal to improve your functional capacity, and your functional capacity improves, you may no longer meet the early intervention requirements anymore.

If you're no longer eligible, we'll help you leave the NDIS making sure you're connected with other services in your community, if you need them. We'll also keep your information, so you can apply again if your situation changes.

Learn more about [leaving the NDIS](#).

What happens if you're not eligible?

If you're not eligible, you can't become an NDIS participant.

We'll try and contact you by phone, or your preferred contact method,⁷⁰ to explain why you're not eligible. We'll give you reasons for our decision and answer any questions you might have.

We'll also send you a letter with our decision, including the reasons you're not eligible and what to do next. Your letter will confirm the date we made the decision.

Even if you're not eligible for the NDIS, your [early childhood partner](#) or [local area coordinator](#) can help you connect to [mainstream and community supports](#).

Your early childhood partner or local area coordinator will use the information you shared to suggest supports in your community. They'll work with you to see how these supports may help you with what is important to you. We also have a list of [other government and community supports](#) you can get, even if you're not eligible for the NDIS.

What if you don't agree with our decision?

If you don't agree with our decision that you're not eligible, you should [contact us](#). We can help explain our decision and what your options or next steps might be.

You can also ask for an internal review.⁷¹ Another staff member, who wasn't involved in the original decision, will then check if we made the right decision. You need to ask for an internal review within **3 months** after receiving the decision.⁷² Learn more about [reviewing our decisions](#).

But you can't ask for an internal review if:

- you withdraw your application because you don't want to apply anymore
- a decision has not been made and we ask you for more information
- you didn't give us information on time and we took this to mean you withdrew your application

- it's been more than 3 months since you received our decision that you're not eligible.

If you don't agree with the internal review decision, you can ask the Administrative Review Tribunal to review it. We call this an external review. You can't ask for an external review until after we make the internal review decision.

You can [contact us](#) to discuss any concerns you may have about the process. You can also [make a complaint](#) if you're not happy with any part of the process.

Can you apply again?

Yes. If we decide you're not eligible, or you're no longer eligible, you can apply again, unless you have requested a review of that decision and are waiting for a decision to be made on the outcome of your review. This includes when you apply for the NDIS or leave the NDIS after your status as a participant has been revoked. Learn more about [leaving the NDIS](#).⁷³

You'll follow the same process to apply as you did the first time. Remember, you need to be younger than 65 on the day you make your new application. And children with developmental delay will need to be younger than 6 on the day they apply.

If you've asked for an internal review of the decision, you can't apply to the NDIS again until we've completed that review.⁷⁴ You can ask us to stop our internal review at any time.⁷⁵

Also, if your review is with the Administrative Review Tribunal after an internal review, you can't apply again until it has made a decision.⁷⁶ You can also ask the Tribunal at any time to withdraw your application. Learn more on the [Administrative Review Tribunal website](#).

Applying to the NDIS – Appendices

What's on this page?

- [List A: Conditions that are likely to meet the disability requirements](#)
- [List B: Conditions that are likely to result in a permanent impairment](#)
- [List C: What if you're receiving disability support in Western Australia?](#)
- [List D: Permanent impairment/Early intervention, under 7 years. No further assessment required](#)
- [When do we make priority eligibility decisions?](#)
- [How do we weigh evidence of disability?](#)

List A: Conditions that are likely to meet the disability requirements

1. **Intellectual disability** diagnosed and assessed as moderate, severe or profound in accordance with current DSM criteria.
2. **Autism** diagnosed by a specialist multi-disciplinary team, paediatrician, psychiatrist or clinical psychologist experienced in the assessment of Pervasive Developmental Disorders and assessed using the current Diagnostic and Statistical Manual of Mental Disorders (DSM-V) diagnostic criteria as having severity of Level 2 (Requiring substantial support) or Level 3 (Requiring very substantial support).
3. **Cerebral palsy** diagnosed and assessed as severe (e.g. assessed as Level 3, 4 or 5 on the Gross Motor Function Classification System - GMFCS).
4. **Genetic conditions** that consistently result in permanent and severe intellectual and physical impairments:
 - Angelman syndrome
 - Coffin-Lowry syndrome in males
 - Cornelia de Lange syndrome
 - Cri du Chat syndrome
 - Edwards syndrome (Trisomy 18 – full form)
 - Epidermolysis Bullosa (severe forms):

- YR
- Autosomal recessive dystrophic epidermolysis bullosa
- Hallopeau-Siemens type
- Herlitz Junctional Epidermolysis Dystrophica
- Lesch-Nyhan syndrome
- Leigh syndrome
- Leukodystrophies:
 - Alexander disease (infantile and neonatal forms)
 - Canavan disease
 - Krabbe disease (globoid cell leukodystrophy) – Infantile form
 - Pelizaeus-Merzbacher Disease (Connatal form)
- Lysosomal storage disorders resulting in severe intellectual and physical impairments:
 - Gaucher disease Types 2 and 3
 - Niemann-Pick disease (Types A and C)
 - Pompe disease
 - Sandhoff disease (infantile form)
 - Schindler disease (Type 1)
 - Tay-Sachs disease (infantile form)
- Mucopolysaccharidoses – the following forms:
 - MPS 1-H (Hurler syndrome)
 - MPS III (San Fillipo syndrome)
 - Osteogenesis Imperfecta (severe forms):
 - Type II - with two or more fractures per year and significant deformities severely limiting ability to perform activities of daily living
- Patau syndrome
- Rett syndrome
- Spinal Muscular Atrophies of the following types:
 - Werdnig-Hoffmann disease (SMA Type 1- Infantile form)
 - Dubowitz disease (SMA Type II – Intermediate form)

- X-linked spinal muscular atrophy
- 5. **Spinal cord injury** or **brain injury** resulting in paraplegia, quadriplegia or tetraplegia.
- 6. **Hemiplegia** where there is severe or total loss of strength and movement in the affected limbs of the body.
- 7. **Permanent blindness** in both eyes, diagnosed and assessed by an ophthalmologist as follows:
 - Corrected visual acuity (extent to which an object can be brought into focus) on the Snellen Scale must be less than or equal to 6/60 in both eyes; or
 - Constriction to within 10 degrees or less of arc of central fixation in the better eye, irrespective of corrected visual acuity (i.e. visual fields are reduced to a measured arc of 10 degrees or less); or
 - A combination of visual defects resulting in the same degree of visual impairment as that occurring in the above points. (An optometrist report is not sufficient for NDIS purposes.)
- 8. **Permanent bilateral hearing loss** > 90 decibels in the better ear (pure tone average of 500Hz, 1000Hz, 2000Hz and 4000Hz).
- 9. **Deafblindness** confirmed by ophthalmologist and audiologist and assessed as resulting in permanent and severe to total impairment of visual function and hearing.
- 10. **Amputation** or congenital absence of 2 limbs – for example, 2 legs, 2 arms, or a leg and an arm (not a leg and a hand, or an arm and a foot).

List B: Conditions that are likely to result in a permanent impairment

Conditions primarily resulting in intellectual or learning impairment

- Intellectual disability
- Pervasive developmental disorders not meeting severity criteria in List A or List C, such as autism
- Asperger syndrome
- Atypical autism
- Childhood autism

Chromosomal abnormalities resulting in permanent impairment and not specified on List A

- Aicardi-Goutières syndrome
- CHARGE syndrome
- Cockayne syndrome Types I and Type II/Cerebro-oculo-facio-skeletal (COFS) syndrome/Pena Shokeir syndrome Type II/Weber-Cockayne syndrome/Neill-Dingwall syndrome)
- Cohen syndrome
- Dandy-Walker syndrome
- DiGeorge syndrome /22q11.2 deletion syndrome/Velocardiofacial syndrome/Shprintzen syndrome/Conotruncal anomaly face syndrome
- Down syndrome/Trisomy 21
- Fragile X syndrome
- Kabuki syndrome
- Menkes disease
- Prader-Willi syndrome
- Seckel syndrome/microcephalic primordial dwarfism/Harper's syndrome/Virchow-Seckel dwarfism
- Smith-Lemli-Optiz syndrome
- Smith-Magenis syndrome
- Spinal muscular atrophy Types III and IV
- Sturge-Weber syndrome
- Trisomy 9
- Tuberous sclerosis
- Turner syndrome
- Williams syndrome
- Wolf-Hirschhorn syndrome

Conditions primarily resulting in neurological impairment

- Alzheimer's dementia
- Creutzfeldt-Jakob disease
- HIV dementia

- Huntington's disease
- Multi-infarct dementia
- Parkinson's disease
- Post-polio syndrome
- Vascular dementia

Systemic atrophies primarily affecting the central nervous system

- Abetalipoproteinaemia
- Adult-onset spinal muscular atrophy/late-onset (SMA type III)
- Fazio-Londe disease/Progressive bulbar palsy of childhood
- Friedrich's ataxia
- Hereditary spastic paraplegia/Infantile-onset ascending hereditary spastic paralysis/L1 syndrome/spastic paraplegias types 2 and 11Huntington's disease/Huntington's chorea
- Louis-Bar syndrome/Ataxia-telangiectasia
- Motor neuron disease/Motor neurone disease/Lou Gehrig's disease /Amyotrophic lateral sclerosis
- Primary lateral sclerosis
- Progressive bulbar palsy
- Spinal muscular atrophy – all types
- Spinocerebellar Ataxia – all types, including Machado-Joseph disease

Extrapyramidal and movement disorders

- Hallervorden-Spatz syndrome/Pantothenate kinase-associated neurodegeneration (PKAN)/neurodegeneration with brain iron accumulation 1 (NBIA 1)
- Parkinson's disease
- Shy-Drager syndrome/Multiple System Atrophy/Striatonigral degeneration (MSA-P)/Sporadic olivopontocerebellar atrophy (MSA-C)
- Steele-Richardson-Olszewski syndrome/Progressive supranuclear ophthalmoplegia
- Stiff-man syndrome/Stiff-person syndrome

Other degenerative diseases of the nervous system

- Alzheimer's disease
- Alpers disease/Grey-matter degeneration/Alpers syndrome/progressive sclerosing poliodystrophy/progressive infantile poliodystrophy
- Lewy body dementia
- Pick's disease

Demyelinating diseases of the central nervous system

- Adrenoleukodystrophy
- Multiple sclerosis
- Schilder's disease/Diffuse myelinoclastic sclerosis – non-remitting

Episodic and paroxysmal disorders

- Brain stem stroke syndrome
- Cerebellar stroke syndrome
- Motor and sensory lacunar syndromes
- Lennox syndrome/Lennox-Gastaut syndrome
- West's syndrome

Polyneuropathies and other disorders of the peripheral nervous system

- Adult Refsum disease
- Charcot-Marie-Tooth disease/Hereditary motor and sensory neuropathy/peroneal muscular atrophy
- Dejerine-Sottas disease/Dejerine-Sottas syndrome/Dejerine-Sottas neuropathy/progressive hypertrophic interstitial polyneuropathy of childhood/onion bulb neuropathy
- Infantile Refsum disease

Other disorders of the nervous system

- Hydrocephalus
- Multiple system atrophy

Conditions resulting in physical impairment

- Amputation
- Congenital absence of limb or part thereof

- Epidermolysis bullosa
- Harlequin type ichthyosis
- Juvenile arthritis / Stills Disease (excluding monocyclic/self-limited Adult Onset Stills disease)
- Rheumatoid arthritis

Diseases of myoneural junction and muscle

- Andersen-Tawil syndrome/Periodic paralysis/myoplegia paroxysmalis familiaris
- Becker muscular dystrophy
- Congenital muscular dystrophy
- Distal muscular dystrophy
- Duchenne muscular dystrophy
- Facioscapulohumeral muscular dystrophy
- Limb-girdle muscular dystrophy
- Mitochondrial myopathy
- Myotonic dystrophy/dystrophia myotonica
- Myotonic muscular dystrophy
- Myotubular myopathy
- Oculopharyngeal muscular dystrophy
- Paramyotonia Congenita
- Thomsen's disease/Congenital myotonia/Becker myotonia

Cerebral palsy and other paralytic syndromes not meeting severity criteria on List A

- Cerebral palsy
- Diplegia
- Hemiplegia
- Monoplegia
- Paraplegia
- Quadriplegia
- Tetraplegia

Conditions resulting in sensory and/or speech impairment

Disorders of the choroid and retina where permanent blindness diagnostic and severity criteria on List A are not met

- Behr's syndrome
- Kearns-Sayre syndrome
- Optic atrophy
- Retinitis pigmentosa
- Retinoschisis (degenerative and hereditary types/juvenile retinoschisis)
- Stargardt disease
- Usher syndrome

Disorders resulting in hearing loss

- Cortical deafness
- Pendred syndrome
- Sensorineural hearing loss
- Stickler syndrome
- Usher syndrome
- Waardenburg syndrome

Conditions resulting in multiple types of impairment

- Aceruloplasminemia
- Addison-Schilder disease/Adrenoleukodystrophy
- Albinism
- Arginosuccinic aciduria
- Aspartylglucosaminuria
- Cerebrotendinous xanthomatosis/cerebral cholesterosis
- Congenital cytomegalovirus infection
- Congenital iodine-deficiency syndrome/cretinism
- Congenital rubella syndrome
- Glycine encephalopathy/non-ketotic hyperglycinaemia

- GM1 gangliosidosis
- Hartnup disease
- Homocystinuria
- Lowe syndrome/Oculocerebrorenal syndrome
- Mannosidosis
- Menkes disease
- Mucopolidosis II/I-cell disease
- Mucopolidosis III/pseudo-Hurler polydystrophy
- Mucopolidosis IV
- Neuronal ceroid lipofuscinosis (NCL)/ Adult type (Kuf's or Parry's disease)/Juvenile (Batten disease)/Late infantile (Jansky-Bielschowsky)
- Niemann-Pick disease
- Pyruvate carboxylase deficiency
- Pyruvate dehydrogenase deficiency
- Sialidosis
- Sulfite oxidase deficiency

The following mucopolysaccharidoses

- Scheie syndrome/MPS 1-H
- Hurler-Scheie syndrome/MPS 1 H-S
- Hunter syndrome/MPS II
- Morquio syndrome/MPS IVA
- Maroteaux-Lamy syndrome/MPS VI
- Sly syndrome/MPS VII

Congenital conditions – cases where malformations cannot be corrected by surgery or other treatment and result in permanent impairment but with variable severity

- Arnold-Chiari Types 2 and 3/Chiari malformation
- Microcephaly
- Fetal alcohol spectrum disorder (FASD)
- Fetal hydantoin syndrome

- Spina bifida
- VATER syndrome /VACTERL association

List C: What if you're receiving disability support in Western Australia?

Please note: the transition of people formerly in Western Australian government disability programs is now complete and List C is no longer in operation.

We have an agreement with the Western Australian government to bring Western Australians onto the NDIS. If you're receiving disability supports in Western Australia, you might already meet most of the eligibility criteria. We'll send you a letter with all the details on how to apply.

The Western Australian government will let us know if you're on a program for faster access to the NDIS. This is called a defined program.

If you're on one of these programs, you need to show us that you:

- are younger than 65 on the day you apply
- live in Australia permanently
- are an Australian citizen or permanent resident.

We'll let you know what evidence you need to give us when you apply.

If you show us you meet the above requirements, we'll decide you're eligible under the **disability requirements**.

Which Western Australian defined programs are eligible for the NDIS?

- WA state-administered National Disability Insurance Scheme
- Supported Community Living
- Community Residential
- Day Options
- Disability Professional Services
- Emergency Accommodation
- Respite
- LAC Coordination
- Recreation

List D: Permanent impairment/Early intervention, under 7 years. No further assessment required.

Synonyms for conditions are also shown (e.g. condition / synonym / synonym).

Conditions primarily resulting in intellectual/learning impairment

Chromosomal abnormalities resulting in permanent impairment

- Global Developmental Delay
- Aicardi syndrome
- Aicardi-Goutières syndrome
- Angelman syndrome
- CHARGE syndrome
- Cockayne syndrome/ Types I and Type II / Cerebro-oculo-facio-skeletal (COFS) syndrome/ Pena Shokeir syndrome Type II / Weber-Cockayne syndrome/ Neill-Dingwall syndrome
- Coffin-Lowry syndrome
- Cohen syndrome
- Cornelia de Lange syndrome
- Cri du Chat syndrome
- Dandy-Walker syndrome
- DiGeorge syndrome/22q11.2 deletion syndrome/Velocardiofacial syndrome/ Shprintzen syndrome/ Conotruncal anomaly face syndrome
- Down syndrome/Trisomy 21
- Edwards syndrome/Trisomy 18
- Fragile X syndrome
- Kabuki syndrome
- Lesch-Nyhan syndrome/Nyhan's syndrome/Kelley-Seegmiller syndrome/Juvenile gout
- Leigh syndrome/Leigh's disease/subacute necrotizing encephalomyelopathy
- Menkes disease
- Patau syndrome/Trisomy 13

- Prader-Willi syndrome
- Rett syndrome
- Seckel syndrome/microcephalic primordial dwarfism/Harper's syndrome/Virchow-Seckel dwarfism
- Smith-Lemli-Optiz syndrome
- Smith-Magenis syndrome
- Sturge-Weber syndrome
- Trisomy 9
- Tuberous sclerosis
- Williams syndrome
- Wolf-Hirschhorn syndrome

Conditions primarily resulting in Neurological impairment

Systemic atrophies primarily affecting the central nervous system

- Friedrich's ataxia
- Hereditary spastic paraplegia/Infantile-onset ascending hereditary spastic paralysis/L1 syndrome/spastic paraplegias types 2 and 11
- Louis-Bar syndrome/Ataxia-telangiectasia
- Niemann-Pick disease (Types A and C)
- Progressive bulbar palsy of childhood/Fazio-Londe disease

The following spinal muscular atrophies

- Spinal muscular atrophy Type I/Werdnig Hoffmann disease/infantile SMA
- Spinal muscular atrophy Type II/Dubowitz disease
- Spinal muscular atrophy Type III Kugelberg-Welander disease/juvenile SMA
- Spinal muscular atrophy lower extremity dominant/SMA-LED
- X-linked spinal muscular atrophy

Extrapyramidal and movement disorders

- Hallervorden-Spatz syndrome/Pantothenate kinase-associated neurodegeneration (PKAN)/neurodegeneration with brain iron accumulation 1 (NBIA 1)

- Alpers disease/Alpers syndrome/Grey-matter degeneration/Progressive sclerosing poliodystrophy/Progressive infantile poliodystrophy
- Demyelinating diseases of the central nervous system
- Adrenoleukodystrophy/X-linked childhood cerebral form
- Alexander disease
- Canavan disease
- Krabbe disease/Globoid cell leukodystrophy
- Pelizaeus-Merzbacher disease

Episodic and paroxysmal disorders

- Lennox-Gastaut syndrome/Lennox syndrome
- West's syndrome

Polyneuropathies and other disorders of the peripheral nervous system

- Dejerine-Sottas disease/Dejerine-Sottas syndrome/Dejerine-Sottas neuropathy/progressive hypertrophic interstitial polyneuropathy of childhood/onion bulb neuropathy
- Infantile Refsum disease

Conditions primarily resulting in physical impairment

- Amputation
- Diamond-Blackfan anaemia
- Epidermolysis bullosa
- Harlequin type ichthyosis
- Hay Wells syndrome/ankyloblepharon/ectodermal dysplasia/clefting [AEC] syndrome
- Joint or limb deformities resulting in impaired mobility
- Juvenile arthritis/Stills Disease
- Osteogenesis imperfecta
- Sjogren Larsson syndrome

Diseases of myoneural junction and muscle

- Congenital muscular dystrophy

- Congenital myotonia/Thomsen's disease/Becker myotonia
- Distal muscular dystrophy
- Duchenne muscular dystrophy
- Emery-Dreifuss muscular dystrophy
- Facioscapulohumeral muscular dystrophy
- Myotubular myopathy
- Oculopharyngeal muscular dystrophy
- Paramyotonia Congenita

Cerebral palsy and other paralytic syndromes

- Cerebral palsy
- Diplegia
- Hemiplegia
- Monoplegia
- Paraplegia
- Quadriplegia
- Tetraplegia

Conditions resulting in sensory and/or speech impairment

- Permanent blindness in both eyes, diagnosed and assessed by an ophthalmologist as follows either:
 - Corrected visual acuity (extent to which an object can be brought into focus) on the Snellen Scale must be less than or equal to 6/60 in both eyes
 - Constriction to within 10 degrees or less of arc of central fixation in the better eye, irrespective of corrected visual acuity (i.e. visual fields are reduced to a measured arc of 10 degrees or less)
 - A combination of visual defects resulting in the same degree of visual impairment as that occurring in the above points.(An optometrist report is not sufficient for NDIS purposes.)
- Deafblindness confirmed by ophthalmologist and audiologist and assessed as resulting in permanent and severe to total impairment of visual function and hearing.

Conditions resulting in multiple types of impairment

- Aceruloplasminemia
- Addison-Schilder disease/Adrenoleukodystrophy
- Albinism
- Arginosuccinic aciduria
- Aspartylglucosaminuria
- Cerebrotendinous xanthomatosis/cerebral cholesterosis
- Congenital cytomegalovirus infection
- Congenital hypothyroidism
- Congenital iodine-deficiency syndrome/cretinism
- Congenital rubella syndrome
- Galactosaemia with long term learning disabilities and neurological impairment
- Glycine encephalopathy/non-ketotic hyperglycinaemia
- GM1 gangliosidosis
- Hartnup disease
- Homocystinuria
- Lowe syndrome/Oculocerebrorenal syndrome
- Mannosidosis
- Menkes disease
- Mucopolidosis II/I-cell disease
- Mucopolidosis III/pseudo-Hurler polydystrophy
- Mucopolidosis IV
- Neuronal ceroid lipofuscinosis
- Niemann-Pick disease
- Phenylketonuria
- Pyruvate carboxylase deficiency
- Pyruvate dehydrogenase deficiency
- Sialidosis

- Sulfite oxidase deficiency

The following mucopolysaccharidoses

- Hurler syndrome/MPS1-H
- Scheie syndrome/MPS 1-S
- Hurler-Scheie syndrome/MPS 1 H-S
- Hunter syndrome/MPS II
- San Fillipo syndrome/MPS III
- Morquio syndrome/MPS IVA
- Maroteaux-Lamy syndrome/MPS VI
- Sly syndrome/MPS VII

The following lysosomal storage disorders

- Gaucher disease Types 2 and 3
- Niemann-Pick disease (Types A and C)
- Pompe disease
- Sandhoff disease (infantile form)
- Schindler disease (Type 1)
- Tay-Sachs disease (infantile form)

Congenital conditions – cases where malformations cannot be corrected by surgery or other treatment and result in permanent impairment

- Chiari malformation/Arnold-Chiari malformation
- Congenital absence of limb(s)
- Congenital hydrocephalus
- Fetal alcohol spectrum disorder (FASD)
- Fetal hydantoin syndrome
- Microcephaly
- Spina bifida
- VATER syndrome (VACTERL association)

When do we make priority eligibility decisions?

If you're in one of the following situations, we'll decide if you're eligible within **2 to 5 business days**.

- Child younger than 7 years with a hearing impairment, either:
 - Identified as Hearing Australia or Early Childhood Partner Priority
 - Identified as 'newly diagnosed'.
- A child is identified as having a developmental delay and is turning 6 years old within 30 days of a valid NDIS application.
- **Immediate risk** to self, others, community or agency where appropriate disability or informal supports are not in place.
- **Unexpected, significant deterioration** of disability related functional capacity where appropriate disability or informal supports are not in place.
- **Rapid deterioration** in functional capacity of a person with one of the following permanent disabilities:
 - Amyotrophic Lateral Sclerosis (ALS or Lou Gehrig's Disease)
 - Brain Cancer
 - Motor Neurone Disease (MND)
 - Progressive Bulbar Palsy (PBP)
 - Primary Lateral Sclerosis (PLS)
 - Progressive Muscular Atrophy (PMA).
- A **terminal illness** and disability.
- **Imminent risk** (within 1 to 14 days) of breakdown of either:
 - Accommodation – risk of homelessness
 - Caring arrangements, including informal supports, due to death, serious illness or injury of informal supports, or significant and unexpected deterioration of disability related functional capacity.
- Appropriate disability supports are not in place and are re-entering the community after a long-term residence or hospital stay (specific release date not required):
 - A person with a **newly acquired, significant disability**, such as spinal cord injury, being discharged from hospital
 - A **younger person living in residential aged care**

- A person being **discharged from an inpatient mental health facility**
- A person due to be **released from correctional facility**.

How do we weigh evidence of disability?

We understand that you may have evidence of your disability from different health professionals at different times. When we're deciding if you're eligible for the NDIS, we look at things like:

- [how old your evidence is](#)
- [who provided your evidence](#).

We weigh evidence based on what we consider best practice, or highest quality. We consider this evidence most strongly when we make a decision. Sometimes, we need evidence from a specific type of health professional. For example, if you have permanent blindness, we need evidence from an ophthalmologist.

What type of evidence should you provide?

We need evidence to help us consider if you meet the disability or early intervention requirements.

For the disability requirements, we need evidence to confirm your permanent impairment and evidence about how this impacts your functional capacity.

For the early intervention requirements, we need evidence to confirm your permanent impairment and evidence that confirms you need early intervention.

It's important to understand the type of evidence that you're providing us. You may have evidence from a doctor or specialist confirming your **permanent impairment** or you may have evidence from an allied health professional or other medical professional that tells us about **impacts to your [functional capacity](#)**. These are different types of evidence which will often be provided by different health professionals based on their qualifications.

How old should your evidence be?

How old should your evidence be to confirm your permanent impairment?

We need evidence from your doctor or specialist to confirm your permanent impairment. You can give us evidence confirming this **from any age**.

How old should your evidence be to confirm your functional capacity?

Generally, we need evidence about how your impairment impacts your functional capacity from **the last 12 months**. This is because your functional capacity may change over time, even if your impairment doesn't.

If you give us more than one type of evidence, we might weigh the newer evidence over the older evidence. If you give us older evidence, we may give this less weighting when we make our decision. In these cases, we'll generally ask for more information. If this isn't provided, we may decide you aren't eligible for the NDIS.

How old should your evidence be to confirm you need early intervention?

We need evidence from your doctor or specialist to confirm your permanent impairment and that you need early intervention.

Generally, we need evidence about that confirms you need early intervention from **the last 12 months**. This is because your functional capacity may change over time – even if your impairment does not.

Who should provide evidence?

We generally prefer evidence that comes from an [appropriate treating professional](#) who:

- is the most **appropriately qualified** person to provide evidence of your primary disability
- has treated you for a significant period of time (at least six months)
- [is registered to practise in Australia or New Zealand](#)
- provides disability evidence (such as a medical report) that is original, genuine and specific to you.

Depending on your situation, you might get your evidence of **permanent impairment** from a different treating professional than your evidence of **functional capacity**.

If you need help to get your evidence together, your [local area coordinator](#) or [early childhood partner](#) can help you.

Who should provide evidence of your permanent impairment?

We generally prefer evidence from your doctor or specialist to confirm your permanent impairment.

Examples of common doctors or specialists include:

- General Practitioner (GP)
- Paediatrician
- Orthopaedic surgeon

- Neurologist
- Psychiatrist.

Who should provide evidence of your functional capacity?

We generally prefer evidence from a doctor, specialist, allied health or other medical professional to confirm how your permanent impairment impacts your functional capacity.

In addition to doctors and specialists, examples of common allied health or other medical professionals include:

- Occupational Therapist
- Speech Pathologist (Therapist)
- Psychologist
- Physiotherapist.

Who should provide evidence that you need for early intervention?

We generally prefer evidence from your doctor or specialist to confirm your permanent impairment.

Whereas a doctor, specialist, allied health or other medical professional can give us evidence to confirm you need early intervention.

In addition to doctors and specialists, examples of common allied health or other medical professionals include:

- Occupational Therapist
- Speech Pathologist (Therapist)
- Psychologist
- Physiotherapist.

For children younger than 6 with developmental delay, an [early childhood partner](#) can give evidence of developmental delay.

Health professionals registered to practise in Australia and New Zealand

We strongly prefer evidence of your disability to come from a registered Australian or New Zealand health professional. Most Australian health professionals are registered with the [Australian Health Practitioner Regulation Agency \(AHPRA\)](#).

We'll still consider evidence from non-Australian or New Zealand health professionals, or unregistered health professionals. However, this evidence will usually be given less weight.

If we can't confirm the registration of your health professional, we'll ask you (and your health professional) for more information in the first instance. If we still can't confirm their registration, we'll give the evidence less weight and will likely decide that you're not eligible for the NDIS.

Reference list

- ¹ NDIS Act s 24(1)(e).
- ² NDIS Act ss 22-23.
- ³ NDIS Act s 24.
- ⁴ NDIS Act s 25.
- ⁵ NDIS Act s 28(1).
- ⁶ NDIS Act s 22.
- ⁷ NDIS Act s 23(1)(b)(i).
- ⁸ NDIS Act s 23(1)(b)(ii).
- ⁹ NDIS Act s 23(1)(b)(iii).
- ¹⁰ NDIS Act s 23(1)(a).
- ¹¹ NDIS Act s 23(2).
- ¹² NDIS Act s 23(2)(f).
- ¹³ NDIS Act s 23(2)(a).
- ¹⁴ NDIS Act s 23(2)(b).
- ¹⁵ NDIS Act s 23(2)(c).
- ¹⁶ NDIS Act s 23(2)(d).
- ¹⁷ NDIS Act s 23(2)(e).
- ¹⁸ NDIS Act s 24.
- ¹⁹ NDIS Act ss 24(1)(a); 25(1)(a).
- ²⁰ NDIS Act ss 24(1)(a); 25(1)(a)(ii).
- ²¹ NDIS Act s 24(1)(b).
- ²² NDIS Act ss 24(3).
- ²³ NDIS (Becoming a Participant) Rules rr 5.4, 6.4.
- ²⁴ NDIS (Becoming a Participant) Rules rr 5.6, 6.6.
- ²⁵ NDIS (Becoming a Participant) Rules rr 5.5, 6.5.
- ²⁶ NDIS Act s 9B(1)(a)(i).
- ²⁷ NDIS Act s 9B(1)(a)(ii).
- ²⁸ NDIS Act s 9B(1)(b).
- ²⁹ NDIS (Becoming a Participant) Rules r 5.8.
- ³⁰ NDIS Act s 24(1)(d).
- ³¹ NDIS Act s 24(1)(e).
- ³² NDIS Act s 24(3).
- ³³ NDIS Act s 24(2).
- ³⁴ NDIS Act ss 25(1)(a)(i)-(ii).
- ³⁵ NDIS Act s 25(1)(a)(i).
- ³⁶ NDIS Act s 25(1)(a)(ii).
- ³⁷ NDIS Act ss 25(1)(a)(i)-(ii).
- ³⁸ NDIS Act s 25(1)(b).
- ³⁹ NDIS Act s 25(1)(c).
- ⁴⁰ NDIS (Becoming a Participant) Rules r 6.9.
- ⁴¹ NDIS Act s 25(1)(d).
- ⁴² NDIS Act s 25(1)(a)(iii).
- ⁴³ NDIS Act ss 9 (definition of 'developmental delay'), 21(1)(c), 25(1)(a)(iii).
- ⁴⁴ NDIS Act ss 9 (definition of 'developmental delay'), 25(1)(a)(iii).

-
- ⁴⁵ NDIS Act ss 9 (definition of 'developmental delay'), 25(1)(a)(iii).
⁴⁶ NDIS Act s 9 (definition of 'developmental delay' para (a)).
⁴⁷ NDIS Act s 9 (definition of 'developmental delay' para (b)).
⁴⁸ NDIS Act s 9 (definition of 'developmental delay' para (c)).
⁴⁹ NDIS Act s 25(1)(d); NDIS (Becoming a Participant) Rules r 6.1.
⁵⁰ NDIS Act s 19(1)(a).
⁵¹ NDIS Act s 19(1)(b).
⁵² NDIS Act s 19(1)(c).
⁵³ NDIS Act s 26(1)(a).
⁵⁴ NDIS Act s 74(1)(a).
⁵⁵ NDIS Act s 74(1)(b).
⁵⁶ NDIS Act s 20(2).
⁵⁷ NDIS Act s 26(1).
⁵⁸ NDIS Act s 26.
⁵⁹ NDIS Act s 26(3).
⁶⁰ NDIS Act s 26(2).
⁶¹ NDIS Act s 21(3).
⁶² NDIS Act s 100(5)(b).
⁶³ NDIS Act s 28(1).
⁶⁴ NDIS Act s 28(2).
⁶⁵ NDIS Act s 28(2)(a).
⁶⁶ NDIS Act s 32BA(1).
⁶⁷ NDIS Act s 32BA(5).
⁶⁸ NDIS Act s 32BA(4).
⁶⁹ NDIS Act s 34(1)(aa).
⁷⁰ NDIS Act s 7(2).
⁷¹ NDIS Act s 100(2).
⁷² NDIS Act s 100(2).
⁷³ NDIS Act s 19(2).
⁷⁴ NDIS Act s 19(2)(c).
⁷⁵ NDIS Act s 102.
⁷⁶ NDIS Act s 19(2)(d).