Life After Stroke

WE’RE ON YOUR SIDE

www.nichs.org.uk
Life after Stroke - the Early Days

Every stroke is **different**. Each person affected by stroke will have different problems and different needs. The way in which you might be **affected** depends on where in the **brain** the stroke happens and the nature of the stroke.

The brain **controls** the way we move, think, see, speak, and eat. **Everything** we do is controlled by different parts of the **brain**.
The left side of the brain controls the right side of the body and the right side controls the left side of the body. So a stroke on the right side of the brain generally causes problems on the left side of the body.

Each side of the brain also has different functions and this will also influence what effect the stroke has, or what symptoms each person experiences.

The left side of the brain normally controls reading, writing, speaking and understanding. So you are more likely to have speech problems if the stroke affects the right side of your body.
Movement, sensation and vision are controlled by both sides of the brain. Therefore these functions can be affected by whichever side of the brain is damaged.

Effects following a stroke

Common effects, that you may experience, following a stroke include:

- **Weakness** or lack of movement in leg and/or arm
- Problems with **balance** and co-ordination
- Trouble **swallowing**
- Problems with **vision**
- Problems thinking or **remembering**
- Trouble **speaking**, understanding, reading or writing
- Shoulder pain or **arm pain or stiffness**
- Feeling **worried** or sad
- Problems controlling your **feelings**
- **Incontinence**
What caused the stroke?

A **stroke** can happen to **anyone** at any age but some people are more at **risk**. There are many **factors** that can increase your **risk** of having a **stroke**. Some of these, such as increasing age, gender and a family history of stroke, cannot be controlled.

**Risk factors** for stroke that we have more control of are generally related to **lifestyle** such as:

- Smoking
- Eating a high fat or high salt diet
- Being physically inactive.

These have an impact on your blood pressure and cholesterol levels. High **blood pressure**, high **cholesterol** and diabetes are medical conditions that we also have some control over in terms of medication and/or **lifestyle** changes. These conditions put you at greater risk of stroke or heart disease. Often people ask **“Why me?”** Some feel vulnerable,
confused, and worried about their health and the future. Some feel sad or disappointed in their bodies and others feel guilty or blame themselves.

For many, the situation seems unfair, causing them to feel angry at themselves and the people they love. Everyone’s reaction is different, but they’re all completely normal and it is important to try to focus your energy on recovering.

**Your Stroke Journey**

If you have a stroke, the stages of your stroke journey will typically be:

- **Going** to hospital
- **Assessment** of your stroke
- Medical interventions
- **Assessment** of the effects of your stroke
- **Rehabilitation**
- **Discharge** from hospital
This booklet will also deal with:

- Physical effects and rehabilitation
- Emotional and mental wellbeing
- Living with a stroke
- Financial and practical support
- The future

**Going to hospital**

Seeing a **doctor** right away is very **important**. The quicker the stroke is diagnosed and treated the better the **chance of recovery**.

Most people who have had a stroke will go to **Accident & Emergency** for initial **care and assessment**. They may then be transferred to a specialised **stroke unit** (not all hospitals have one) or a general rehabilitation ward. Sometimes with **less severe strokes** people can remain at **home** and come to hospital for tests as an **outpatient**.
Assessment of your stroke

Care in the early stages concentrates on assessment of your stroke and prevention of further damage and complications and may include:

- finding out what the problem is, the type of stroke you have had, the area of damage and how serious it is
- having a brain scan as soon as possible and certainly within 24 hours to help with diagnosis
- receiving drugs to help dissolve the blood clot (thrombolysis)

Not all Hospital Stroke Units offer thrombolysis, and even in units that do, only a small proportion of patients are suitable for this treatment. To be effective, the quicker the drug is given the better. Ideally the drug should be given within three hours of the very first symptom.
• a referral to a brain surgeon for assessment if you have had certain types of brain haemorrhage
• preventing or treating medical problems and complications.

Common tests initially include:

• Blood pressure test
• A chest x-ray (to exclude other health conditions)
• A heart tracing (to check for other forms of heart disease)
• Blood tests (checking cholesterol, blood sugar and clotting)
• A brain scan (to detect affected areas)

Other more detailed tests may be carried out at a later date to confirm what caused the stroke.

Medical Interventions

During your first few days in hospital, care concentrates on assessment and medical interventions to reduce
your risk of further complications or of having another stroke.

Your **treatment** may involve:

- **Blood thinning** drugs (stops clots forming and prevents another stroke)
- Lowering high **blood pressure**
- Carotid **surgery**
- Tube **feeding** if you cannot swallow safely
- **Controlling** risk factors such as diabetes and high blood cholesterol

Once your medical condition is stable many other **healthcare professionals** will be involved in your care. In hospital they will focus on assessing how your stroke has affected your **abilities**. They will work out an individual **care** and **rehabilitation** plan for you. In the early stages you may have **difficulty** concentrating or **retaining information**. It will help to have a **family member** or **friend present** when you are talking to your doctor or other healthcare professionals as they can help to remind you what was discussed as...
well as ask questions you might not think of. It can be useful to write information down so it can be read later.

Assessment of the effects of your stroke
Assessment of memory, understanding and communication

You will be assessed for any problems in your thinking, your memory or concentration, and how aware you are of your surroundings as well as your ability to communicate.

Stroke can affect your ability to speak, write and understand what is being said. This is known as aphasia and can affect people in different ways such as:

- difficulty in speaking or producing any sounds at all
- problems in thinking of the right words to speak or write
- trouble understanding speech or writing or slurring of speech
The effects of your stroke on speech, reading and writing will be fully assessed by a speech and language therapist.

Assessment of Swallowing

After a stroke some people have problems swallowing fluids or food (called dysphagia). Your swallowing should be assessed as soon as possible. Assessment involves a simple test; for example you may be asked to try to swallow a small amount of water. If you have problems swallowing, a speech and language therapist or specialist nurse will carry out further tests and advise you and the staff looking after you on techniques for safe swallowing and on the consistency of food and fluids you should have.

Assessment of Movement

Stroke can cause weakness or paralysis on one side of your body and problems with balance or
co-ordination. You may also experience changes in sensations like tingling, numbness or feelings of hot and cold.

Your ability to move will be assessed as soon as possible after admission. The degree of physiotherapy and occupational therapy you receive will depend on how much movement you have lost as a result of your stroke and on how active you were before you had it.

Assessment of Continence

A stroke can lead to loss of bladder control (urinary incontinence), bowel control (faecal incontinence) or both. Most patients regain continence in a few weeks as their body recovers from the effects of their stroke.

A continence nurse can help with the problems that some people have controlling their bladder and bowel immediately after a stroke. For
example, a **catheter** tube **might** be needed for a short time to help drain the bladder.

**Care on the ward**

The nursing staff will oversee your **care** on the ward depending on how much you are able to do for yourself. You may need **help** getting in and out of bed or getting washed and dressed.

It is important to avoid prolonged time in **bed**, even in the early days to:

- Prevent your **limbs** becoming stiff and sore
- Help regain your sense of **balance**, movement and posture
- Reduce the risk of **blood clots** in the legs or chest infections
During the initial recovery period you will feel **physically tired** and it is extremely important to try to get sufficient **rest** and not to overdo things. The physical effects of the stroke are often worse when tired so **activity** should be **paced** throughout the day.

**Rehabilitation**

Once you are medically **stable** and have been assessed it is likely that the stroke team will work out an individual **care** and **rehabilitation** plan for you.

**Rehabilitation** is the process of overcoming or **learning** to cope with the effects of the stroke. It is about becoming as **independent** as possible by:

- Relearning skills
- Learning new skills
- Adapting to your limitations
Rehabilitation begins in hospital but continues after you go home.

The physiotherapist will support you in becoming more mobile and regaining muscle strength and control. This will help with balance problems, paralysis and muscle weakness.

**Early treatment** will include guided movements of your limbs and making sure you are correctly positioned in bed or your chair. They will develop exercises to improve movement and to stop weak limbs becoming stiff. As you improve your physiotherapist will show you more complicated exercises that encourage your whole body to work together.

The occupational therapist (OT) will teach you new ways to undertake everyday tasks. **Weakness** in your arm or hand or problems with **memory** or thinking can make it difficult to do everyday tasks such as getting washed and dressed or making a meal. The occupational therapist can advise you on new ways
to do things, and **equipment** that might help you at home like handrails or ramps.

The **speech and language therapist** will give you and your family advice on how best to communicate together. They will also monitor the **safety** of your **swallow** and advise what consistency of food and drinks are safe for you to eat and drink on discharge. A **dietician** can also advise on appropriate foods to eat to ensure you get a **balanced** nutritional **diet**.

**How long will I be in hospital?**

The length of **time** you will be in **hospital** depends on a number of factors, in particular, the nature of your stroke. Some people are in hospital for a few days, others a few weeks and occasionally for some, longer than this.
What happens when I am discharged?

Most people are keen to go home and this is a very important motivator for recovery. But it can also be daunting for the person who has had the stroke, their carers and family.

All the members of the hospital team will be involved in preparing for your discharge and will make arrangements for further treatment after discharge if required.

Doctor

The hospital doctor will give you or your family a short note containing the diagnosis of the stroke and a list of the medication to be continued after discharge. This will be followed up by a full and detailed letter to your GP to make sure that they have all the necessary clinical information required for your ongoing care.

Usually the hospital will give several days supply of
the necessary medication. Your GP will then issue a health service prescription and in the long-term, supplies can be obtained at your local chemist.

The doctor will also ensure that you are on the appropriate medication to provide as much protection as possible against another stroke.

You may be given an appointment to attend the hospital clinic, generally between one and three months after discharge. This provides a useful opportunity to discuss any outstanding issues.

**Nurse**

The nursing staff play a central role in co-ordinating your discharge plan, to ensure that all your needs are met.
They will assess your **nursing needs** in hospital and will be able to advise on ongoing issues, such as **continence, skin care** and the need for additional **equipment**. They will contact nursing colleagues in the **community** if appropriate.

**Physiotherapist**

The physiotherapist will advise on what help you need with **mobility** after **discharge** and ensure that appropriate arrangements are in place. They will also provide relevant **practical advice** to your family or carer.

**Occupational therapist**

If you are being discharged home, the **occupational therapist** will take into account the layout of your house, and may recommend specialised **equipment** or adjustments needed to help you **live** in your own **home**.
Speech and language therapist

If you still have **problems** with **communication** or **swallowing** a community speech and language therapist may also see you after you get home.

Social worker

The **social worker** will support you through the **discharge process** and provide **emotional support** to you and your **family**. They will ensure that the necessary arrangements are put in place in time for discharge. They will make contact with **community services**, and organisations like Northern Ireland Chest Heart & Stroke, where appropriate.
What will life be like after I get out of hospital?

It’s difficult to give advice that applies to everybody. Many people need to make adjustments after a stroke. It is important that you receive the necessary support from your family, friends and healthcare professionals.
We will now talk about:

- Community Stroke Team
- Physical effects and rehabilitation
- Emotional and mental wellbeing
- Living with a stroke
- Financial and practical support
- The future

**Community Stroke Team**

A team of *health professionals* will be involved in your rehabilitation. The team might include, doctors, nurses, physiotherapists, occupational therapists, speech and language therapists, dieticians and social workers depending on your needs and the resources in your area.

The *team* will work closely together with you and your family. It is important that you ask the team any *questions* you have or ask for an explanation of
anything you do not understand.

**Northern Ireland Chest Heart & Stroke** also have a team of Stroke Family Support Co-ordinators who can offer **support** to the carers and families of people who have had a **stroke**. Stroke Family Support offers advice and information as well as practical and emotional support. Contact Northern Ireland Chest Heart & Stroke on **028 9032 0184** for more information.

**Physical effects and rehabilitation**

Some people do not require further **rehabilitation** after their initial treatment. If you require ongoing rehabilitation you may receive it in your **home**, or as an outpatient at the **hospital**.

The length of time your **rehabilitation** continues will depend on the problems you have and the progress you are making as well as the **services** that are
available in your area.

The aim of rehabilitation is to become as independent as possible within the limitations of your stroke.

You can help yourself with the recovery process by:

- Remaining **positive**
- Being **patient** and not over exerting yourself
- Doing **exercises** given to you by health professionals
- Following **medical advice**
- Making **adjustments** and finding new ways of doing things
- **Talking** to your friends and family
- **Accepting** help and support

**Working towards recovery**

**Physical recovery** from stroke depends on the effort you make, and so it is important that you are **positive** and that you practice your rehabilitation exercises
and activities in-between sessions with the health professionals.

**Tiredness** is a common problem for some time after stroke and your problems can appear worse when you are tired. Try to avoid having too many **visitors** especially soon after discharge, and take a rest everyday. Do your **exercises** or activities during short sessions throughout the day. Congratulate yourself on reaching targets set by the health professionals or by yourself and then concentrate on the next realistic target. Setting smaller, more achievable **goals** is better for your motivation than setting goals which are harder to achieve and will take much longer for you to see **progress**. However, don’t feel you need to force yourself to go on if you are tired.

The time will come when your **structured rehabilitation** will end but this does not mean that you will not
make any further progress. It is important for you to continue to set yourself realistic goals and to work towards them using the skills that you have learned during rehabilitation.

Emotional and mental wellbeing

Every stroke is different and everyone will have their own experience of stroke but most people will ask “Why me?” Hospital tests will have identified, in most cases, the cause of your stroke. However, because stroke is often sudden it can cause feelings of shock as well as denial, anger, guilt and loss for the life you have led.

Almost like grief there is a journey of emotions and acceptance that you will go through after your stroke. Initially denial protects you from being overwhelmed by the changes in your life. Over a longer time denial blocks your progress both physically and emotionally.
Anger is a normal response once you have accepted that you have had a stroke and that life may have changed. Some people feel angry at themselves, some at God, at a loved one or at the doctors and nurses. Often anger is directed at those closest to you and this can cause problems.

Feelings of guilt are often reported after stroke. People can feel guilty about what caused the stroke, about the worry they are causing to loved ones and about the effect on family life, for example not being able to work. It is important not to bottle up all these feelings. Talking to your family and friends might not be easy but it will help, and remember that they have been affected by your stroke too, and talking will also help them as well.

Northern Ireland Chest Heart & Stroke have Stroke Family Support Co-ordinators who can help you by talking things through with you and your family.
Contact Northern Ireland Chest Heart & Stroke on 028 9032 0184 for more information.

Some people find it helpful if they don’t compare life the way they are now with the way they were before the stroke happened. It is important to focus on what you can do now. It may be necessary to be flexible and try new ways of doing the things that are important to you.

**Emotional Control**

Changes in your emotions or behaviour may be due to damage to the part of the brain that controls emotions. Your emotional reactions can be exaggerated and sometimes inappropriate to the situation. Some people experience sudden unprovoked tearfulness, inappropriate laughter or outbursts of anger.

These reactions are often distressing for both you
and your family or friends. Understanding that they are another symptom of stroke can help you all cope with the situation.

This is usually a temporary situation and in time you will have more control and feel more like yourself. However not all emotional changes are due to the effects of brain injury. Some may be due to depression or anxiety.

_I am feeling a bit down, is this normal?_

Sometimes depression can be the direct result of the damage caused by the stroke. Other times it may be as a result of the emotional difficulty involved in coming to terms with having a stroke.

_Feeling down_ some of the time is natural after a stroke but if you are feeling low most of the time you could be depressed. Depression can develop soon after the stroke or months afterwards.

Common signs of depression include:
• Feeling **sad** or “empty” most of the time
• **Fatigue** or not wanting to get up in the morning
• Changes in **sleep** patterns, difficulty sleeping or early morning waking
• Changes in **appetite**
• **Mood** swings
• Loss of **motivation**
• Lack of **interest** in people or activities
• **Crying** all the time
• Feelings of **despair** or thoughts of suicide

**Depression** is very **common** and doctors are very familiar with it and how to treat it. Your **doctor** may suggest you try taking **antidepressant medication**.

Many people have been helped through a **difficult time** in their lives by taking antidepressant medication until they have **come to terms** with their **new life**. Your **GP** can advise you on whether **medication** is appropriate to you or suggest other ways of coping.
It can also **help to talk** to other people who **understand** what you are going through. **Northern Ireland Chest Heart & Stroke** provide **Stroke Schemes** for people who have had strokes. **Contact** Northern Ireland Chest Heart & Stroke on **028 9032 0184** for more information.

**I am feeling a bit anxious, is this normal?**

**Anxiety** is a feeling of fear or nerves. **Anxiety** after stroke is usually related to feelings of fear:

- **Fear** of having another stroke
- **Fear** of not being able to **cope** in life
- **Fear** of the **unknown**

Feelings of anxiety can cause **physical** and **psychological** symptoms.

Physical symptoms of anxiety:
- A racing heart or irregular heartbeat (palpitations)
• Excessive sweating
• Dizziness
• Over breathing (hyperventilation)
• Frequent urination, or feeling a need to go to the toilet
• Muscle aches and tension
• Nausea
• Dry mouth

Psychological symptoms of anxiety:
• On-going worry or fear that doesn’t seem to go away.
• Feeling ‘on edge’ or ‘wanting to run away’
• Restlessness
• Irritability
• Poor concentration / easily distracted
• Feelings of panic or panic attacks
• Avoiding situations that may trigger anxious feelings
It is common for people to suffer both anxiety and depression at the same time. There are medications and therapies that can help with both problems. If you start to feel anxious it may help to try some breathing exercises. Often just concentrating on your breathing will take your mind off your immediate worries and it also helps reduce your heart rate and some of the other physical signs of anxiety.

Tips for breathing control

- Settle yourself in a relaxed position with your back supported
- Place your hands between your lower ribs and navel. Your shoulders should be relaxed
- **Breathe** in through your nose; you should feel your tummy move out as you breathe in, 1..2..3..4
- Breathe out gently through your lips, your tummy will move in as you breathe out, 1..2..3..4
- Concentrate on the **rhythm** of your breathing, feel your muscles relaxing and let all other thoughts drift away
• Practise this breathing control regularly so that you are able to use it when you really need it.

If you do feel depressed or anxious talk to your loved ones about your fears and anxieties. It can help. Talking will also help them as they will have their own concerns about you.

It can also help to talk to other people who understand what you are going through. Northern Ireland Chest Heart & Stroke provides support through Stroke Family Support and Stroke Schemes for people who have had strokes. Contact Northern Ireland Chest Heart & Stroke on 028 9032 0184 for more information.
Other Issues

There are other mental health difficulties that people experience after a stroke such as Agoraphobia (fear of open spaces or going outside), health anxiety and social anxiety. If you feel any emotional distress it is worthwhile talking to your GP. If you are have feelings of despair or thoughts of suicide contact Lifeline on 0808 808 8000.

Can I drive after my stroke?

A stroke is a condition which could affect your driving ability and may need to be reported to the DVA (formerly the DVLNI).

Everyone will have to stop driving for one month after a stroke. Whether or not you can return to driving after a month depends on how the stroke affected you:

• if there are no lasting effects you may be advised
that you can return to driving at the end of the first month
• if the stroke has affected your vision this may mean that you will be advised not to drive, but further tests may be required
• if the stroke has affected your arm or leg function this may also affect your ability to drive after the initial month, and you will be advised not to drive while further assessments may be needed

Lasting limb disability does not necessarily prevent you from driving in the long term. Adaptations to a vehicle and/or restrictions to automatic vehicles may enable you to drive. In all cases seek medical advice from your GP about your ability to drive. It is also important to note that it is your responsibility to inform the DVA. You should also inform your car insurance provider.

Resuming a social life

Some people find social situations difficult after a stroke. You might avoid people. You might fear
questions about your stroke or that people will feel sorry for you.

You might feel **self-conscious** about any physical or communication difficulties that you have. You may **worry** about how you will manage when you are out of the house.

It may seem easier to stay at home but this will lead to **isolation**. At first try to go out for short periods of time. You will confront the situations you are fearful of gradually. **Testing** your **ability** to cope will **increase** your **confidence** over time. You will be relying on other people to understand your situation and to co-operate but you may be pleasantly surprised at how accepting and helpful most people are. Don’t be disheartened before you try. Over time you will become more comfortable and will enjoy getting out and socialising again.
Resuming Sport and Leisure

Your physiotherapist will be able to give you advice on how to maintain the physical level you have reached during rehabilitation.

One way of maintaining exercise and activity is to become involved in sport and leisure activities. They are also a good way to socialise with other people.

You may be able to do sports you enjoyed before your stroke if you are determined enough. Many sports can be adapted to become accessible to people with different abilities. For example, many people play one-handed golf or get involved in wheelchair sports. All ability sports such as Boccia and New Age kurling are also becoming popular.

Your local council will be able to provide information about swimming clubs or exercise classes which cater for people with disabilities.
Can I go on holiday after my stroke?

If your stroke has left you with any disability you are not advised to fly for about six weeks and if you do plan to fly you should consult your doctor.

After three months, as long as you feel well enough and your GP has no objections, there is no reason why you should not take a holiday. Holidays can be a great way to recharge the batteries and relax. So start as you mean to go on, be organised and:

- Think about where you are going and if it will be easily accessible for you
- Allow yourself plenty of time to get to the location
- Don’t be carrying heavy pieces of luggage around
- Contact the airline or holiday company to discuss any mobility aids or additional help you may need at least 48 hours in advance.
- Plan your holiday so that you do and see all the
things you want to without having to rush around

• Take some time to relax through the days to avoid being overtired.

You may want to book with a travel agent who can provide specialist advice or book accommodation equipped for your stay. If the flight is over two hours, it is best to take a walk along the aisle at regular intervals so that you reduce the risk of a DVT (blood clot) in the legs. Also you could exercise your calf muscles for several minutes every half an hour to help circulation in the legs. It is also advisable to avoid alcohol and caffeine (tea, coffee and cola). Drink plenty of water to avoid dehydration, which is especially common during night flights.

Remember to take enough medication which will last you through the holiday. If you are flying, carry it in your hand luggage so that it is accessible and unlikely to get lost. It is also a good idea to take a list of your medications with you.
It is also important to make sure you have adequate travel insurance. Northern Ireland Chest Heart and Stroke has a list of insurers. Call 028 9032 0184 or go to www.nichs.org.uk/travelinsurance. If you are travelling within the EU it can be useful to also carry the European Health Insurance Card (EHIC) which entitles holders to free medical treatment in some European countries or to claim back some medical costs. You can apply for the card through the NHS Business Service Authority. See www.ehic.org.uk. The EHIC will not cover all costs so it is still important to get adequate travel insurance.

**Will I be able to return to work?**

Returning to work may be important for personal and financial reasons and so it can be a motivating factor for recovery. However, going back to work can be difficult after a stroke if you are experiencing tiredness, memory problems, difficulty concentrating and any physical disability.
If going back to work is right for you it is important not to return too soon. It is important for you to talk to your employers about their expectations and your own expectations. Some changes may need to be made by your employers and by you, for example, changes to your working pattern, to help you return to and carry out your job.

If you are unable to go back to the same job and want to continue to work, then you can consider other options such as changing jobs, working part-time, volunteering, retraining or returning to education.

More advice can be obtained from the Disability Employment Officer at your local Training and
Employment Agency.

Volunteering can be a great way of building up confidence within a working environment. It can also be very fulfilling. Many Northern Ireland Chest Heart & Stroke clients go on to volunteer within our organisation.

**Can I have sex after a stroke?**

Sexual difficulties after stroke can develop for many different reasons. The emotional physical and psychological impact of stroke can create problems. These issues can be short-term and relatively easy to overcome, but for others, they are more serious and longer lasting.

Disabilities, such as weakness or paralysis, may cause problems due to physical limitations. Some medications are also known to have side effects that may affect your sex life.

Having a stroke often affects the way you see yourself,
resulting in a loss of self-confidence, lowered self-esteem and lack of interest in sex.

The first step in dealing with any sexual problems is to talk about them with your partner. It may also help to discuss the issue with your doctor as they may be able to resolve some of the physical difficulties or those caused by medication.

It is also important to remember that simply being close to someone is essential to help a person feel loved, special and truly a partner in the relationship.

**HRT after stroke**

Current research shows that if you have had a stroke you should not take HRT (Hormone Replacement Therapy) as it will slightly increase your risk of having another stroke or developing other health problems. If you have been taking HRT before having a stroke, your doctor will recommend that you stop taking it immediately.
What benefits or allowances am I entitled to?

The Social Security Agency assesses and pays benefits to people in Northern Ireland. The benefits system is complex and it changes frequently so it is important to get advice about what you are entitled to and how to fill in the necessary forms.

Information and advice can be obtained from the Social Security Agency. Any Citizens Advice Bureau or Independent Advice Centre can give you advice about what you are entitled to.

The table opposite will give you an indication of which benefit(s) you may be entitled to.
<table>
<thead>
<tr>
<th>Age</th>
<th>Employment situation</th>
<th>Benefit</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Under 16 years old</strong></td>
<td>n/a</td>
<td><strong>Disability Living Allowance (DLA)</strong></td>
</tr>
<tr>
<td><strong>16-64 years old</strong></td>
<td><strong>Employed but off work sick</strong></td>
<td>You may be able to claim <strong>Statutory Sick Pay</strong> (SSP) from your employer for up to 28 weeks depending on how long you have been employed, your age, and how much you earn. When your entitlement to SSP ends you may be able to claim <strong>Employment and Support Allowance</strong>.</td>
</tr>
<tr>
<td><strong>16-64 years old</strong></td>
<td>Employed and back at work</td>
<td>If you go back to work, you may be entitled to an increase in <strong>Working Tax Credit</strong>.</td>
</tr>
<tr>
<td>Age Group</td>
<td>Eligibility</td>
<td>Benefits</td>
</tr>
<tr>
<td>---------------------------</td>
<td>---------------------------</td>
<td>----------------------------------------------</td>
</tr>
<tr>
<td>16-64 years old</td>
<td>Unable to return to work</td>
<td>Disability Living Allowance (DLA).</td>
</tr>
<tr>
<td>Over 65 years old</td>
<td>Retired</td>
<td>Attendance Allowance, which is for people who need assistance with their personal care and/or supervision.</td>
</tr>
<tr>
<td>Over 16 years old (no upper age limit)</td>
<td>Carer (at least 35 hours per week)</td>
<td>If the person you are caring for receives DLA (middle or higher rate) or Attendance Allowance, you may be eligible for Carer’s Allowance. You do not have to be a relative of or live with the person you are caring for.</td>
</tr>
</tbody>
</table>
What other support can I receive?

Community support services to help you manage at home are usually arranged through Social Services within the Department of Health, Social Services and Public Safety.

You may already have been referred to a community social worker. If you haven’t seen a social worker your GP can refer you to Social Services.

You will have to have your needs assessed by Social Services before they will provide services for you. This is called a community care assessment.

The assessment should take into account:

- Your wishes as the person being assessed
- Whether you have any particular physical difficulties which affect your ability to manage everyday tasks including personal care
- Whether you have any particular housing needs
- What sources of help you have access to, such
as carers, family or nearby friends, and their willingness to continue providing care

- What needs the people who provide care may have

Once you have been assessed, the social worker can tell you about what services are available and whether you have to pay for them. Services can vary from area to area but might include:

- A home help or care assistant
- Delivery of meals to the home
- A place at a day centre or within a community group

Whilst community care assessments and carer’s assessments are free, Social Services can charge
for some community care services. The regulations about which community care services must be paid for, and how much can be charged, are complicated. The Citizens Advice Bureau can give you independent advice on accessing community care services.

Recognise that the stroke will change your life but that this need not prevent you from having new hopes and goals for the future.

**Will I have another stroke?**

The risk of having a stroke increases if you have already had a stroke or TIA. Your risk also depends on the type of stroke you experienced and the treatment you had at the time. If you had an operation it may have fixed the problem causing your stroke. Likewise, your doctors may have given you medications to treat underlying conditions and to reduce the risk of further stroke.

There are risk factors that may increase your risk of stroke that you cannot control. Age, gender and your
family’s medical history all play a role in the risk of another stroke.

There are a lot of risk factors that you CAN control.

You should:

- Do regular exercise
- Choose a healthy diet
- Stop smoking
- Drink alcohol in moderation
- Manage stress

It is difficult to make lifestyle changes but is definitely worth the effort. Please contact NICHS for further information on health promotion and stroke prevention.

The Future

Stroke is traumatic. It brings about a lot of life changes all at once. It will take time to mentally and physically adjust to life after stroke. It is never possible to predict the future. This uncertainty can cause a sense of fear.

You may never get back to exactly how you were before your stroke but you may continue to make
improvements overtime. You will learn to adjust and to find new ways of doing things that are important to you. You will have **good** days and **bad** days.

Stroke does not prevent you from having new **hopes** and **goals** for the future. Many stroke survivors have discovered new talents and interests and made new friends as part of their life journey after stroke.

**What can NICHS do to help?**

Northern Ireland Chest Heart & Stroke (NICHS) offers a wide range of **stroke schemes** for people who have had strokes or TIAs. Services provided include:

- **Information** and **education**
- **Self-management** programmes
- Practical and emotional **support**
- Health promotion and stroke **prevention**
- **Advocacy**
- **Liaison** with other health professionals
- Support schemes

NICHS have a team of **Stroke Family Support Co-ordinators** across Northern Ireland who can offer...
information and support to the carers and families of people who have had a stroke.

NICHS services can be accessed at any stage of your life after stroke. We can arrange home visits if you have difficulty travelling to our venues.

Contact Northern Ireland Chest Heart & Stroke for more information on 028 9032 0184 or mail@nichs.org.uk.