What is primary progressive aphasia (PPA)?
This leaflet is about primary progressive aphasia (PPA), a rare form of dementia. It’s for anyone who may be worried about themselves or somebody else.

The information here does not replace any advice that doctors, pharmacists or nurses may give you. It provides background information that we hope you will find helpful.

It was written in May 2017 and will be reviewed in May 2019. Please get in touch if you’d like a version with references or in a different format.

This information booklet was produced in partnership with Rare Dementia Support.
What is dementia?

The term dementia is used to describe a group of symptoms, including memory loss, confusion, language and communication problems, mood changes and difficulty with day-to-day tasks.

The symptoms of dementia get worse over time, so that a person will need more and more help with everyday life.

There are many causes of dementia. These include Alzheimer’s disease, vascular dementia, dementia with Lewy bodies and frontotemporal dementia.

What is primary progressive aphasia?

Primary progressive aphasia (PPA) is a condition caused by damage to parts of the brain that control our personality, emotions, language and behaviour. In most cases, this damage is caused by frontotemporal dementia. Most people who develop PPA will be in their 50s and 60s.

The term PPA covers three separate conditions. In all three, people’s speech and language is usually affected first, but in different ways:

**Semantic dementia**
Over time, people forget the meaning of words as well as objects.

**Progressive non-fluent aphasia**
People find it harder to speak and are more likely to make mistakes in how they say words or sentences.

**Logopenic aphasia**
People often pause while speaking as they try to find the right word.
## Semantic dementia

People with semantic dementia gradually find it harder to remember the meaning of words. For example, they may:

- **Have trouble using the right word** – often saying another word or using a vague term like ‘thing’. This may start with names and other words they don’t use very often. As time passes, people will also struggle with more common words and often ask what words mean.

- **Forget what everyday objects are for** or find it hard to recognise them.

- **Talk about things at great length** and in a vague or roundabout manner.

- **Find it hard to understand** what other people are saying.

- **Have problems reading and spelling**.

In the later stages, people tend to say less. Changes in behaviour and personality are more common in semantic dementia than the other types of PPA. For example, people may develop obsessions or a sweet tooth, or act in ways that may seem strange to others.

Later they may find it difficult to recognise people they know, things around the house or familiar sounds. This makes it harder to get on with day-to-day life.

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## Progressive non-fluent aphasia

This condition affects how a person produces speech. For example, they may:

- **Have trouble producing words**, although they know what they want to say. Speaking may take a lot of effort and words may not come out right.

- **Find that words come out in the wrong order** or are missed out altogether.

- **Stutter or speak more slowly** or hesitantly and be hard to understand.

- **Struggle to use the right grammar** when speaking or writing letters or emails.

If you use sentences that are long and complex, someone with this form of dementia may find it hard to understand you. Over time, they may develop other signs including:

- **Problems with reading, writing and spelling**.

- **Trouble understanding some words**.

- **Changes in behaviour**.

- **Trouble with swallowing**.

- **Problems making decisions or plans**.

Some people show signs similar to Parkinson’s disease such as shaking, being unsteady on their feet or having trouble using their hands.
Logopenic aphasia
This form of PPA was identified more recently. People with logopenic aphasia are generally able to speak and understand others but have trouble finding the words they want to use. For example, they may:

Have trouble remembering the right word. The person may pause as they try to find the word they want.

Speak more slowly and hesitantly, and find it hard to say words correctly.

As time passes, people can have more problems with their memory and thinking. For example, someone may forget conversations or appointments, become lost, find it hard to work out how much change they need for a bus fare or struggle to use household appliances, like washing machines.

With all three types of PPA, problems get worse over time. It becomes harder for people to say what they need and move about on their own. Dementia is different for everyone who has it, and the speed of change can vary widely. However, someone with PPA will need increasing care and support until the end of their life.
Diagnosis

It is important to get the right diagnosis so that the right help can be given. If you are worried about your health or someone else’s, you should talk to your doctor.

If your doctor suspects PPA or another form of dementia, they may send you to a specialist clinic.

Here, a doctor or nurse will run through more detailed tests with you. These may include:

• Questions about your symptoms and medical history.
• Speaking to your partner or someone close to you about the problems you are having.
• A physical check-up.
• Assessments of your thinking and language skills, and memory.

You may also have blood tests, an MRI brain scan or other specialist tests.

Together all of these will help a doctor to work out what is causing the speech, language and other problems.

PPA is a rare type of dementia, not often seen by doctors. It may take longer than usual to get the right diagnosis.

When a diagnosis is made, many people will want to make plans for the future. This includes care options, legal and money matters. Find out about help and support for families affected by dementia on page 14.
Treatments

There are currently no medicines specifically for PPA, but there are treatments that might help with some of the symptoms.

For example, speech therapy can help with language problems. Your doctor can refer you to a Speech and Language Therapist, who can also help those having trouble swallowing food.

People with all forms of dementia may be offered antidepressants, or non-drug treatments like cognitive behavioural therapy (CBT). Ask your doctor for more information.

What causes PPA?

PPA is caused by loss of brain cells in the front and sides of the brain, called the frontal and temporal lobes. The reasons for this loss are not yet clear.

We know that there is an unusual build-up of certain proteins inside brain cells. These proteins include TDP-43 and tau but it’s a complicated picture. Researchers are working hard to find out why this happens and how it damages brain cells.

Most people with logopenic aphasia have the same kind of damage in their brains as people with Alzheimer’s disease. This includes the build-up of a protein called amyloid in the brain. Therefore, in most cases logopenic aphasia is an unusual form of early-onset Alzheimer’s.

In a small number of people semantic dementia or primary non-fluent aphasia may be caused by a faulty gene that can be passed down in families. The genes involved are called MAPT, progranulin (or GRN) and C9ORF72. These genes are also associated with other forms of frontotemporal dementia. For more information, ask for our ‘Genes and dementia’ booklet.
A diagnosis of PPA will affect people in different ways, and they may need time to get used to it. With the right information and support, people can carry on with regular aspects of their lives for some time. Talking to other people in the same situation can help.

The PPA Support Group holds several meetings a year, helping people living with PPA to meet and talk to others. Visit the PPA Support Group website at www.raredementiasupport.org/ppa. email contact@raredementiasupport.org or call 07341 776 317.

YoungDementia UK is a charity offering support for people under 65 with any form of dementia, and their families. Their website, www.youngdementiauk.org, has a range of resources, including information about local services. They also have a Facebook group: www.facebook.com/YoungDementiaUK. Call 01865 794311.

Admiral Nursing Direct offers practical and emotional support to anyone affected by dementia, including advice on managing the symptoms. Call 0800 888 6678.

Alzheimer’s Society provides information and help for people with all forms of dementia, and can tell you about local support groups and Dementia Cafés in your area. Call 0300 222 1122.

The PSP Association website, www.pspassociation.org.uk, has some helpful resources and an online forum. PSP (progressive supranuclear palsy) is a different condition, but some of the symptoms are similar, including speech and swallowing problems, and occasionally PSP can begin with PPA.

You can also talk to your doctor or nurse for advice on caring for someone with PPA.
Help and support

With all types of PPA, speech or language is affected first.

When talking to someone with PPA, there are things you can do to help them:
• Be patient and understanding.
• Find a quiet place to talk.
• Don’t try to finish a person’s words unless they ask you to help.
• Speak loudly and clearly.
• Check that you have understood what they mean.
• One-to-one conversations may work better than talking in groups.
• If the person starts to find talking hard, take a break so they can rest.
• They may find it easier to write things down or use an electronic aid than to talk.

If someone is finding eating and swallowing hard, these ideas may help:
• Try not to have foods like thick pieces of meat, or cook them slowly so they are easier to chew and swallow.
• If food is getting stuck in the throat, avoid dry and crumbly foods. Add sauces to meals and make sure people have plenty of water to drink with their food.
• If eating becomes slower, a plate-warmer will help to keep food warm and nicer to eat.
Help and support

With time, someone with PPA will need more help with day-to-day life. It is important to think about safety at home and any changes you may need to make. In those of working age, PPA may make working life more difficult. It is helpful to talk to close family, early on, about options such as lasting power of attorney, and later, home care and care homes. Driving safety is also important. You should let the DVLA know if you or a family member is diagnosed with any form of dementia.

For organisations offering help and support, ask for a copy of our booklet 'Caring for someone with dementia: organisations that can help' or visit our website www.alzheimersresearchuk.org

Research

Alzheimer’s Research UK has funded over £12 million of pioneering research into different forms of frontotemporal dementia.

Our research is gaining new insights into the proteins that build up in the brain and how these damage nerve cells as well as taking these discoveries into drug discovery programmes to develop new treatments.

We believe that dementia can only be defeated through research. Thanks to the generosity of our supporters, we hope to fund many more pioneering studies into the diseases that cause dementia.

Find out more

If you have questions about dementia research or want to find out more about how to get involved in studies, contact our Dementia Research Infoline on 0300 111 5 111 or email infoline@alzheimersresearchuk.org

The Infoline operates 9.00-5.00pm Monday to Friday. Calls cost no more than national rate calls to 01 or 02 numbers and should be included in any free call packages.
We are the UK’s leading research charity aiming to defeat dementia.

We welcome your comments to help us produce the best information for you. You can let us know what you think about this booklet by contacting us using the details below.

**Contact us**
Alzheimer’s Research UK
3 Riverside, Granta Park, Cambridge CB21 6AD

**Supporter care**
T: 0300 111 5555
E: enquiries@alzheimersresearchuk.org

**Dementia Research Infoline**
T: 0300 111 5 111
E: infoline@alzheimersresearchuk.org

**www.alzheimersresearchuk.org**

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### Send me more information

For free information, simply complete this slip. You can drop it straight in a post box or put it in an envelope labelled with the freepost address overleaf. Alternatively, phone us on **0300 111 5555**.

**I would like to know more about**

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<th>Topic</th>
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<td>Treatments for dementia (SCIHTFD)</td>
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<td>Genes and dementia (SCIIGENE)</td>
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<td>Caring for someone with dementia: organisations that can help (SCIICARE)</td>
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<td>The latest dementia research (SMTHINK)</td>
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**Name**

**Address**

We’d really like to keep you updated with the latest research developments, how your support is making a difference, and fundraising activities that you can get involved in. Your details are always held securely, but if you’d rather not hear from us please tick the relevant box:

- No information by post [ ]
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