Positive language
An Alzheimer’s Society guide to talking about dementia
What is positive language and why is it important?

There are currently 850,000 people living with dementia in the UK, each with their own unique personalities and life stories. They have the right to be treated with respect, and not stigmatised or treated differently to others because of their dementia.

The way we talk about dementia has a direct effect on how people living with the condition feel. It can also have a profound effect on society. After all, the words we use affect the way we think, and the way we think affects how we behave. The language we all use has the power to impact the lives of millions of people around the country.

For those living with dementia, using words or phrases that label, belittle or depersonalise people can have a big impact on them and their family and friends. It changes the way they feel about themselves, shaping their mood, self-esteem, and feelings of happiness or depression. It can also change the way other people think about dementia, and increase the likelihood of a person with dementia experiencing stigma or discrimination.

On the other hand, using positive language – words and phrases that empower people, treat them with dignity, and respect them as individuals – can change how society views and treats people with dementia. It can make people feel valued and included, and contribute to changing the landscape of dementia.

This is our guide to using positive language. It has been informed by what we have been told by people with dementia, their carers and all those affected by the condition. By following these guidelines you can help improve people’s ability to live well with dementia, and bring about lasting social change.

This guide includes how we should talk about:
- dementia
- people living with dementia
- the effects of dementia
- people caring for a person with dementia
- Cultural awareness and dementia
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The main rules and principles
There are some important basic principles to follow when writing about dementia.

**Do use the following**

- Always use language that recognises dementia is not the defining aspect of a person’s life. **See the person, not the dementia.**

- Always think about how a person with dementia would feel about your choice of words. Respect people’s wishes about how they want to be talked about, whether they are present or not and whether you think they can understand or not. If you are unsure of what words to use, or how to talk about someone, ask them what they prefer.

- When you are telling someone’s story, use **their own words as much as possible**, as long as it is appropriate. Dementia affects everyone differently, and everyone will express their experiences of dementia in their own way. If you can, ask the person directly.

- Use language that **focuses on what people can do, rather than what they can’t**, or that shows someone that you are with them but not doing something for them. This can help people stay positive and retain feelings of self-worth. For example, use phrases like ‘supporting a person to eat’, rather than ‘feeding’.

**Don’t use the following**

- **Avoid labelling people**, or using terms that place the emphasis on the condition or behaviour, rather than the person. For example, don’t label someone – a person may have dementia, but they are not a ‘dementia sufferer’; they may be experiencing changes in behaviour, but they are not ‘a shouter’.

- **Don’t reinforce stereotypes or myths about dementia.** It is important to know the facts and avoid assumptions. For example, it is not a normal part of ageing, nor is memory loss the only symptom or always present in the early stages.

- **Do not use abbreviations to describe people.** For example, do not use ‘PWD’ (person/people with dementia) or ‘YPWD’ (younger person/people with dementia).
2 Talking about dementia
Dementia is not a disease. It is a condition that is caused by diseases:

- Dementia is the name for a group of symptoms that may include memory loss and difficulties with thinking, problem-solving or language, as well as changes in perception, mood or behaviour.

- Dementia is caused by diseases that affect the brain. The most common of these is Alzheimer’s disease. However, there are many causes of dementia and they affect people differently.

- Dementia is not just about memory loss, and many people with dementia will not have memory loss. While some people may prefer to speak of their ‘memory problems’ rather than their dementia, we should not use the two synonymously as this can exclude people with other types of dementia, or who have different symptoms.

Young-onset dementia

If someone develops dementia before the age of 65, they have young-onset dementia, or alternatively working age dementia or early-onset dementia (note that this doesn’t mean the early stages of dementia). Never use senile or pre-senile dementia.

Senile dementia is an outdated term that was used when it was thought that symptoms associated with dementia, such as memory loss or difficulties with thinking, were just a normal part of ageing, rather than being caused by diseases that affect the brain.

Use the following ways of describing dementia:

- dementia
- Alzheimer’s disease and other types of dementia
- a form of dementia
- a type of dementia
- a condition
- symptoms of dementia.

Don’t use the following:

- dementing
- demented
- affliction
- senile dementia
- senility.
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Talking about people with dementia
People with dementia are individuals with a history, personality, relationships and desires. Their diagnosis is not the defining aspect of their life, nor does it mean that their life is over. We should use language that reflects this.

Use the following when talking about a person with dementia:
- A person (or people) with dementia
- A person (or people) living with dementia
- A person (or people) with a diagnosis of dementia (if appropriate – remember not all people with dementia get a diagnosis).

Using these terms helps to maintain the person’s dignity. They put the person first and don’t make assumptions or pass judgment on people because of their dementia.

However, try to avoid using ‘person with dementia’ or ‘person living with dementia’ too often, as it can become almost another label. Vary the language used – for example, ‘someone who has dementia’, or, ‘Mrs Jones, who has Alzheimer’s disease’.

Some carers may describe themselves as ‘living with dementia’, because they feel like they live with the condition by living with a person with it. We talk more about this on page 15.

Never use the following:
- sufferer, suffer from, afflicted, victim or any other words that disempower people with dementia by making them seem passive, childlike, or worthy of pity
- senile, pre-senile or demented
- slang expressions that are derogatory – for example, ‘not all there’, ‘delightfully dotty’, ‘away with the fairies’, ‘doolally’, ‘a few cups short of a tea set’, ‘a few pennies short of a pound’
- patient – don’t describe someone as a ‘dementia patient’. The term ‘patient’ should only be used specifically if a person is in hospital, or in relation to seeing their GP or other healthcare professional.

Younger people with dementia

There are currently more than 42,000 people in the UK under 65 who have dementia. You should describe someone with dementia who is under 65 as a younger person with dementia, or a person with young-onset dementia.
Talking about the effects of dementia
Positive language – How to talk about the effects of dementia

Symptoms

Dementia affects every person differently. This depends partly on the type of dementia the person has, but also on their personality and experiences, and how other people respond to them. The nature and severity of symptoms are likely to change over time.

When talking about symptoms, describe the change itself – for example, ‘memory loss’, or ‘problems finding the right word’. Never label someone as a result of their symptoms. For example, if someone is experiencing changes in behaviour that cause them to walk about or shout out, they are not a ‘wanderer’ or a ‘shouter’. Labels such as these dehumanise the person and place the focus on the behaviour rather than what the person may be trying to achieve or express. This can also make it more difficult to identify what is behind a person’s behaviour, which makes it harder to address it.

A lot of people dislike the term ‘wandering’ if a person walks about because it implies movement without purpose. The person may have a clear purpose or motivation – they may be looking for somebody or something, or walking as a way of calming down – but they may not be able to express it easily. Use ‘walking’ or ‘walking about’ instead.

Challenging behaviours

Changes in a person’s behaviour or emotional state are common in dementia. However this will often be a sign of a person’s physical or emotional need or a feeling that they are not able to communicate. This could be for a number of reasons, for example anxiety, frustration, pain, boredom, loneliness or confusion.

The terms ‘challenging behaviours’ and ‘behaviours that challenge’ are used to refer to a specific subset of changes in behaviour. These include aggression, agitation, shouting or walking about. This is because they are a challenge for the person experiencing them, as well as someone caring for them.

They are often the result of needs or desires that are not being met or cannot be communicated. There is a reason the person is behaving that way – they are not simply ‘being difficult’. For example, if a person is shouting or is agitated, it may be because they are in pain but cannot communicate it. A third ‘challenge’ is to work out what is causing the person to behave this way.

As with all phrases, be careful not to repeat ‘challenging behaviours’ or ‘behaviours that challenge’ too often. Try using variants such as ‘behaviours that the person might find difficult’, or ‘distressed behaviours’. Be careful not to use ‘difficult behaviours’ as this can imply that the person experiencing the behaviour is being difficult.

Think carefully before using ‘behavioural and psychological symptoms of dementia’. It is very medicalised and is used to refer to a broad range of things, from apathy, depression and anxiety to psychoses (hallucinations, delusions) and behaviours that challenge (such as agitation or aggression). Not all of these, however, are inevitable symptoms of dementia. There may be a variety of other reasons for them.
Use the following terms to describe changes in a person’s behaviour or emotional state:

- changes in behaviour
- challenging behaviour
- behaviours that challenge
- distressed behaviours
- symptoms of distress
- behaviour that is out of character
- expressions of needs or desires that are not being met
- behavioural and psychological symptoms of dementia (in a clinical context).

Don’t use the following:

- difficult behaviours
- ‘being difficult’.
The impact of dementia

Because every person’s experience of dementia is different, its impact on people’s lives will be varied and not always negative. Use language that reflects this. While it is important to be truthful and realistic about the impact of dementia, we don’t need to use words that are negative, disempowering, pessimistic or frightening.

Use the following when describing the impact of dementia:

- challenging
- life-changing
- stressful

Don’t use the following:

- hopeless
- unbearable
- impossible
- tragic.

It is OK to use these words if this is how someone has described their own personal experience of dementia, though you should always make it clear when you’re quoting someone directly. However, we shouldn’t assume these things or put words in someone else’s mouth.
Talking about people caring for a person with dementia
The term ‘carer’ generally refers to people who provide unpaid care, such as a partner, family member or friend. It is helpful to use different words for professional or paid carers, such as ‘care worker’ or ‘care professional’.

Not everyone will like to be referred to, or even see themselves as a carer, although this language is widely used in the health and social care system (for example, a carer’s assessment, Carer’s allowance). If possible, ask the person how they feel about this term. It may change over time.

You should also think before assuming that someone is a carer just because they are the partner of a person with dementia. Many people, particularly in the early stages of dementia, don’t need any care yet. They may find it patronising and presumptuous if their partner is automatically referred to as their carer.

‘Living with’ and ‘affected by’ dementia

Most people with dementia say that they prefer the term ‘living with dementia’ to be used for a person who has dementia. Although family, carers and friends are all affected, it is usually not helpful to say that they are also ‘living with dementia’.

The term ‘affected by dementia’ is a good way to refer to a much larger group of people - it can include not only people living with the condition, but their carers, friends, family, or anyone else who is close to them or provides support. For example, ‘when writing this guide we consulted widely with people affected by dementia’ – because we spoke to people with dementia, but also these other groups as well.
Don’t use the following:

- **Burden** – this can be used in different ways, and context is key:
  - Everyone has a different experience as a carer, and using terms such as ‘burden’ in relation to a person’s caring role assumes that role is entirely negative or always difficult. Of course, a carer may talk about the difficulties they experience as a result of providing care, and they may use this word themselves, but we should not make assumptions. Avoid ‘burden’ unless a carer uses the word themselves - use ‘impact’ or ‘effect’ instead.
  - We should never refer to a person themselves as being a burden. This dehumanises someone, and makes them out to be nothing more than a drain on time and resources rather than a human being.
  - In the broader context of health and social care policy, we may talk about ‘the burden of unpaid care’ – this refers to the cost (in terms of time, money and effort) that unpaid carers are having to shoulder on behalf of society. In this broad context, when we aren’t referring to specific people but to issues of policy, the term ‘burden’ is acceptable.

- **Sitting service or sitter** – this implies that the person with dementia is a passive recipient of care, and also has connotations of babysitting.

- **Feeding, dressing or toileting** – again, these terms imply that the person is passive and does not participate in the activities described. When talking about specific tasks, use phrases like ‘supporting (or helping) the person to eat’.

**Use the following when talking about those caring for a person with dementia:**

- supporting a person with dementia
- caring for a person with dementia
Glossary of preferred terms
The following is a list of DOs and DON’Ts – however this list is for guidance only. It is not exhaustive and does not cover everything. Also, some of these words are OK if you are reporting how someone affected by dementia describes their own experience of the condition.

<table>
<thead>
<tr>
<th>Topic</th>
<th>Preferred terms</th>
<th>Do not use</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dementia</td>
<td>dementia (a condition)</td>
<td>dementia (a disease)</td>
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<tr>
<td></td>
<td>Alzheimer’s disease and other types of dementia</td>
<td>dementing illness</td>
</tr>
<tr>
<td></td>
<td>a form of dementia</td>
<td>demented</td>
</tr>
<tr>
<td></td>
<td>a type of dementia</td>
<td>affliction</td>
</tr>
<tr>
<td></td>
<td>symptoms of dementia</td>
<td>senile dementia</td>
</tr>
<tr>
<td></td>
<td>young-onset dementia</td>
<td>senility</td>
</tr>
<tr>
<td>People living with dementia</td>
<td>a person (or people) with dementia</td>
<td>sufferer</td>
</tr>
<tr>
<td></td>
<td>a person (or people) living with dementia</td>
<td>suffer from</td>
</tr>
<tr>
<td></td>
<td>a person (or people) with a diagnosis of dementia</td>
<td>afflicted</td>
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<td></td>
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<td>victim</td>
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<td></td>
<td></td>
<td>demented</td>
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<td></td>
<td></td>
<td>patient (when used outside the medical context)</td>
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<td></td>
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<td>not all there</td>
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<td></td>
<td></td>
<td>lost their mind</td>
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<td></td>
<td></td>
<td>derogatory slang expressions – for example, delightfully dotty, doolally, away with the fairies</td>
</tr>
<tr>
<td>People with dementia under 65</td>
<td>young-onset dementia</td>
<td>pre-senile dementia</td>
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<td></td>
<td>working age dementia</td>
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<td></td>
<td>early-onset dementia</td>
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### Positive language – Glossary of preferred terms

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<tr>
<th>Topic</th>
<th>Preferred terms</th>
<th>Do not use</th>
</tr>
</thead>
<tbody>
<tr>
<td>The symptoms of dementia</td>
<td>Describe the symptom itself – for example: memory loss, difficulty communicating, changes in behaviour, expressions of unmet need, challenging behaviour, behaviours that challenge, behavioural and psychological symptoms (in a clinical context)</td>
<td>wanderer, Shouter, wandering, difficult behaviours, being difficult</td>
</tr>
<tr>
<td>The impact of dementia</td>
<td>challenging, disabling, life changing, stressful</td>
<td>hopeless, unbearable, impossible, tragic</td>
</tr>
<tr>
<td>People caring for a person with dementia</td>
<td>caring for a person with dementia, supporting a person with dementia</td>
<td>living with dementia</td>
</tr>
<tr>
<td>The act of providing care to a person with dementia</td>
<td>caring, supporting/helping the person to eat/dress/go to the toilet</td>
<td>sitting service, sitter, feeding, dressing, toileting, burden (for an individual)</td>
</tr>
</tbody>
</table>
The word ‘dementia’ may mean other things to people from different cultural backgrounds or who speak other languages. It can be seen as offensive or disrespectful. Although the understanding of dementia is generally growing, and the stigma around it generally reducing, be aware and sensitive to the fact that this change is happening differently in each community.

In some languages there is no equivalent word for dementia, and the closest word sometimes has implications of ‘madness’. In some cultures dementia is associated with mental illness, or seen as a natural part of aging. Words and phrases that are appropriate in one community may be offensive, ambiguous or meaningless in another. It is important to be aware of the cultural background of the person, family or community so that you can use the most appropriate language. Ask for guidance from the relevant community where possible.
For further information

Dementia words matter: Guidelines on language about dementia (The Dementia Engagement and Empowerment Project [DEEP], 2014). Available at: http://dementiavoices.org.uk/2014/10/dementia-words-matter/

Acknowledgments

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