Living with dementia
Your essential guide

What’s inside

Reliable expert advice
Written by our specialist dementia nurses

Practical tips
Hear from people with lived experience of dementia

Heard on the Helpline
Answers to the most commonly asked dementia questions
If you love someone living with dementia, you’re living with it too.

It’s always there. But so are we. See how our specialist nurses can help your family live better with dementia.

dementiauk.org/Together
#ILiveWithDementia
I’m so pleased you’re reading this guide.

At Dementia UK, we know that a dementia diagnosis doesn’t only affect the person with the condition; it can be life-changing for close family and friends too, as you face new and complex challenges together.

**But you’re not alone.**

Every day, our dementia specialist Admiral Nurses help people just like you. We give expert practical advice – from managing behaviour changes to unlocking financial help – and we provide emotional support, helping you look after yourself while you look after the person with dementia.

During my many years working as an Admiral Nurse, I’ve often seen carers putting their own needs to one side. **But if you don’t take care of yourself, it’s harder to care for someone with dementia.** Think of it like putting on your own oxygen mask on a plane before you can help anyone else.

That’s why we’ve created this guide. Inside, you’ll find expert, reliable advice to help you feel more confident, reassured and prepared.

This isn’t an A-Z of everything you’ll ever need to know about dementia – you can find all that on the Dementia UK website. But each section features the most common questions we hear on our Helpline, practical solutions from people who’ve experienced what you’re going through, and expert tips from Admiral Nurses.

I hope you find it useful. And if you need to talk to one of our Admiral Nurses, please get in touch. You’ll find all of our contact information at the end of this guide. Dementia is always there, but so are we.

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**Dr Hilda Hayo**  
Chief Admiral Nurse and CEO, Dementia UK
What’s inside

This guide is divided into eight sections. In each one, you’ll find helpful information for looking after someone with dementia.

Along the way, you’ll read advice and tips from our dementia specialist Admiral Nurses (you can meet some of them over the page), as well as people who’ve experienced what you’re going through.

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✓ You don’t need to read this guide from front to back – skip to the sections that sound most useful right now.

✓ Why not print out this guide at home? You can stick the most relevant pages on the fridge as a reminder.

✓ We’ve included some space (page 40) to jot down notes – and you’ll find useful contacts at the back.

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Meet the Admiral Nurses

If you love someone with dementia, we’re here for you. This guide aims to make life easier by bringing together the most important information about living with dementia in one place.

To help, we’ve asked four of our dementia specialist Admiral Nurses to guide you through each section – highlighting the most useful tips and advice along the way. Let’s meet them.

Meet Admiral Nurse Vic

“I became an Admiral Nurse in 2007. Three members of my own family have had dementia, and I’ve seen first-hand the impact the condition has had.

“Like you, I long for the day when we can say we have a cure for dementia, or that we can prevent it.

“Until that day, we need to make sure no family has to face dementia alone by providing specialist support and advice when they need it. I know every family living with dementia is doing whatever they need to do – day in, day out. I feel privileged to share my time and knowledge to help in any way I can to make those days a little easier.”

Meet Chief Admiral Nurse and Dementia UK CEO Hilda

“After developing and leading specialist dementia teams and services throughout the NHS, I became the Chief Admiral Nurse and Chief Executive Officer for Dementia UK in 2013. I’m so proud of the positive difference our charity makes to people living with dementia. But this guide is for you: the carers, friends, family, and loved ones. You’re also living with dementia and need our support.

“I hope this guide will help you and your family navigate your way through the challenges of dementia, find support and plan for the future.”
All about Admiral Nurses

What is an Admiral Nurse?
Admiral Nurses are dementia specialist nurses – every one of us is a qualified nurse with a professional specialism in dementia. We provide life-changing support for families affected by all forms of dementia. And we’re here to help when you need it most, with expert knowledge across all aspects of dementia.

What do we do?
We support everyone facing dementia: the person with the diagnosis and the people caring for them. This could be by helping the person with dementia live more independently or giving their family members advice on how to plan for the future. Admiral Nurses are here for everyone, making sure people living with dementia and their carers have all the support they need, when they need it.

Where can you find us?
Admiral Nurses always have the time to listen and offer expert advice. You can call our free Helpline seven days a week, book an appointment through our face to face or virtual clinics, or ask your GP, social worker or hospital staff if there is a community Admiral Nurse service in your area. We’re here for you, for as long as you need us.

How we can help
We don’t provide hands-on nursing care, so an Admiral Nurse won’t visit your home to care for your loved one. But we do put families in contact with other care professionals, offer practical and emotional support during all stages of the dementia journey, help coordinate the different services supporting the person with dementia, and much more.

Meet Admiral Nurse Gary
“I’ve been working for Dementia UK for just over two years, where I cover both the Helpline and our virtual clinics. I’ve always been interested in dementia and helping families – I wanted to be an Admiral Nurse for 20 years, so I was really excited when the perfect role came up in my area.
“I know a lot of family members become a carer for someone with dementia without any preparation or training. That’s why it’s vital for them to have access to the right information, advice and expert support. I hope this guide will give you just that, helping you adjust to your new role and look after your own wellbeing.”

Meet Admiral Nurse Kerry
“I feel incredibly privileged to be an Admiral Nurse, supporting families living with dementia.
“I joined Dementia UK two years ago as the Consultant Admiral Nurse for Acute Services. Recently, I’ve moved into a new role as Consultant Admiral Nurse for Frailty.
“When you’re looking after someone around the clock, you might find it difficult to think about your own needs. You can only care for the person with dementia if you look after your own health and wellbeing. Reach out for help and remember we’re here to support you too.”
How to get a diagnosis of dementia is one of the most common questions people ask our dementia specialist Admiral Nurses.

What you need to know

• **signs:** common symptoms of dementia include difficulties with short-term memory, problem-solving, communication, motivation and being able to carry out everyday tasks

• **impact:** the impact of a diagnosis can affect the person with dementia and everyone around them: family, friends, colleagues and employers. Every situation is unique, and how you feel may change over time, too

• **patience:** getting a dementia diagnosis may take time. Waiting lists for memory assessments differ around the UK and you could see a number of consultants before diagnosis. It can also take longer if the person is younger

• **process:** you can ask for a second opinion if your GP won’t refer your loved one to a specialist, but they don’t have to agree

• **you’re not alone:** our Admiral Nurses can answer your questions about getting a diagnosis, the impact it may have, what happens next, and where to go for more help

I wish I’d known...

“People living with dementia can mask how they’re feeling. Just as children can cover up their difficulties, many adults with dementia know they’re struggling but can blag their way through even the toughest situations.

“My mum hid her dementia really well – she didn’t want people to think she was weak or that she didn’t understand them. That was her superpower, but it was also her downfall because nobody knew when Mum did need help.”

Lisa, who cared for her mum

Heard on the Helpline

“I think my partner has dementia but I’m not sure what to do.”

If someone you love is showing signs of dementia, it’s important for them to see a GP as soon as possible. Getting diagnosed means everyone can start planning for the future. Symptoms of dementia can also be similar to those caused by an infection, diabetes, a vitamin B12 deficiency, depression or anxiety. A GP can rule these out or refer the person for assessment at a memory clinic.

“My dad is refusing to see the GP – how can we make him go?”

This is a natural reaction; sometimes people with dementia symptoms are frightened about being diagnosed, or they may not be aware that they’re having problems. Explain that you’ve noticed a few changes, and that they could be caused by something else that’s easily treated. A GP can work out what’s behind those changes, and getting a prompt diagnosis means your loved one can get the right support sooner.

“I’m struggling to cope with Mum’s dementia diagnosis.”

Even if you suspect dementia, it can still be a shock when it’s diagnosed. Having it confirmed makes it ‘real’ and can lead to new, difficult questions – who do you tell? How will they take it? What does it mean for the future? It’s normal for some people to feel relieved after getting a diagnosis, as it explains what’s been happening – try not to feel guilty if that sounds like you.
Quick Q&A

Chief Admiral Nurse Hilda answers some common diagnosis questions.

Q. Do I really need to get an official diagnosis?
A. Getting a diagnosis may feel daunting, but it can help a person access benefits and support, plan for the future and protect them from discrimination if they’re working (dementia is recognised as a disability).

For some types of dementia, the person may also be prescribed medications that might help with symptoms.

Q. What happens during a dementia assessment?
A. A GP will carry out a physical examination and order blood tests to check for other causes, then ask some simple questions to test memory and brain function.

If they have concerns, the GP may refer the person to a memory clinic, psychiatrist or a neurologist for further assessment.

A specialist will then do more in-depth tests, such as counting backwards from 20, remembering and recalling an address, or drawing a clock. They might also carry out a brain scan.

Life hack

“Once you’ve been referred to a memory clinic, you can contact them directly – many families don’t realise this and end up waiting much longer for an assessment.

“If you do call, check that they’ve received the referral from your GP; sometimes letters go missing. And ask to be considered for any cancellations, especially if you can get there at short notice.”

Admiral Nurse Gary
top tips

What are the best things to do when waiting for an assessment? Admiral Nurse Vic has these useful suggestions:

1. **Keep a symptom diary**, noting down any changes. Show this to the specialist so they can clearly see any issues or developments.

2. **Try a cognitive stimulation therapy activity book.** These are packed with games and puzzles to help memory, comprehension and attention.

3. **Encourage the person to keep doing activities they enjoy**, such as swimming, baking, shopping or listening to music. Spending time outside and reconnecting with nature can also help boost their mood.

4. **If your loved one has a job**, they should consider telling their boss/HR department that they’re being assessed for dementia. This can help their employer prepare in advance for any support or adjustments the person might need to keep working.

5. **Start a life story book.** Include your loved one’s interests, likes and dislikes, a bit about their childhood or working life. This will give health professionals a better understanding of their needs in future.

**NHS this way**

Click here for information on GP services and referrals for specialist care.
What helped me most

“Coming to terms with my diagnosis – once I did that, it helped me find my new normal. My life doesn’t revolve around my diagnosis, but it has helped me learn to live life to the fullest. I think it’s incredibly important to keep doing the things you enjoy, and to get the most out of life.”

Jude (right), who has young onset dementia
When someone is diagnosed with dementia, it’s natural to worry about how it will affect their day-to-day life. But there are steps you can take to help them live well with dementia.

**What you need to know**

- **understanding**: living well with a diagnosis starts with understanding. The right information can explain what dementia is and how it’s likely to affect your loved one, you and your whole family. Be mindful that this can change over time, but you can reach out to our Admiral Nurses at any point.

- **planning together**: make sure any lifestyle changes are right for the person with dementia. Try to involve them in important discussions before you make those changes.

- **start small**: introduce any adjustments gradually. Too many new changes can create confusion, so start with little things that will be the most useful.

- **expert advice**: you may need extra support accessing local health or care services. We’re here to help you navigate your way through the system.

- **staying positive**: there’s so much uncertainty around life with dementia: will you need to change anything at home? How will it affect your loved one’s mental health? What impact could it have on the rest of your family? Although it will affect things, we’re here to help you know that there’s life after a dementia diagnosis.

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**I wish I’d known...**

When someone who drives is diagnosed with dementia, they’re legally required to tell the DVLA (DVA in Northern Ireland). This doesn’t necessarily mean they’ll have to give up driving immediately, but they may need a medical and/or driving assessment to renew their licence.

Visit DVLA England, Scotland and Wales [here](#), and DVA Northern Ireland [here](#).

“Ian had always been a passionate driver and I wanted him to continue driving for as long as possible. As soon as he was given the go ahead by doctors, we took a family road trip to Snowdonia. Ian was on top of the world – quite literally, after we all climbed Snowdon together! It was a wonderful lasting memory for us.”

Marion, whose husband Ian had young onset Alzheimer’s disease

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**What helped me most**

“Lester started refusing to change his clothes – he’d wear them for days on end and wouldn’t let me wash them. Our Admiral Nurse Sam suggested I get lots of clothes that were exactly the same, all in one colour. Then I’d quickly swap them round when Lester was in bed.”

Diana, who cares for husband Lester
Quick Q&A
Admiral Nurse Kerry answers some quick questions about living with dementia.

**Q. What’s the best dementia equipment to buy?**
**A.** This depends on what your loved one really needs, but a large clock that tells the time, day and date is very useful. This prevents the person with dementia having to think about it and feeling anxious if they don’t know the answer.

**Q. Why are they not eating or drinking?**
**A.** Dementia means some people don’t recognise when they’re hungry or thirsty, or they may forget that they haven’t eaten. Encourage the person with dementia to eat well by involving them in preparing food, cooking with different flavours and colours to stimulate appetite, making mealtimes a social occasion and including plenty of foods that contain fluids, like watermelon and yoghurt. Give them a drink with every meal and serve it in their favourite cup or glass. You could also make an appointment with their dentist to rule out any dental problems.

**“Your life isn’t over”**
“My advice to anyone with dementia is, ‘Your life isn’t over’. Do what makes you happy, whether that’s seeing friends, walking, or whatever you enjoy. I’ve taken up painting; my school art teacher was wrong about my creative abilities! Now I have a new network of friends that I regularly meet up with. For me, dementia has been the start of a new chapter – and it’s not a bad chapter.”

George, who has mixed dementia

**Heard on the Helpline**

“Where can I find specialist dementia technology?”
There are a number of websites that sell assistive technology for people living with dementia, such as personal alarms, automatic pill dispensers, light timers or GPS trackers – **find out more on our website**. But does your loved one really need all the gadgets? Many smartphone apps do the same job, while some simple home adjustments can be just as helpful – and cheaper!

“Are we eligible for any grants to make home improvements?”
There is support available to help a person with dementia live in their own home for longer. Talk to their GP or social services to arrange needs and/or home assessment to work out what might help. You can also ask for a free visit from an occupational therapist who can provide practical support and advice. You may be able to get a grant from your local council to then help pay for any improvements – our Admiral Nurses can help you find more information.

“Dad’s having trouble sleeping. Can anything help?”
Disturbed sleep and difficulties around bedtime are very common. Keep a sleep diary to see if you can spot any causes and track changes in the person’s sleeping patterns. Make sure they’re comfortable (not too hot or cold), have been to the toilet, and their bedroom is calm and quiet. If they’re reluctant to go to bed, wearing your own night clothes or putting on a dressing gown can show them it’s time for sleep.

“Your life isn’t over”
“My advice to anyone with dementia is, ‘Your life isn’t over’. Do what makes you happy, whether that’s seeing friends, walking, or whatever you enjoy. I’ve taken up painting; my school art teacher was wrong about my creative abilities! Now I have a new network of friends that I regularly meet up with. For me, dementia has been the start of a new chapter – and it’s not a bad chapter.”

George, who has mixed dementia

**How to help the person with dementia maintain a healthy diet**

Click here to watch seven top tips

[Image of healthy food]
Bright ideas for around your home

Try these simple – yet effective – changes to help make everyday life a little easier.

**Front of house**
- Make sure the outside of your home is recognisable and avoid making any changes. Get a large door number and avoid changing the colour of the front door.

**Bedroom**
- A wardrobe full of clothes can feel overwhelming – switch to a simple clothes rail. Use pictures to show what’s in each drawer, like socks, underwear or jumpers.
- Mirrors can be confusing, as the person with dementia may not recognise themselves or think someone else is in their home. Cover them up when not in use.

**Bathroom**
- Swap the toilet seat and towels for brighter colours, as contrasting colours can be seen more clearly.
- Put labels or pictures on the door – like a sink, a toilet or a bath – to help the person with dementia navigate around their home more easily.

**Kitchen**
- Make things easily accessible – remove cupboard doors or put labels on the front showing what’s inside. Pictures may be more helpful than words, such as a can of beans on the food cupboard.
- Keep any frequently used items on the kitchen worktop, like cups, teabags and spoons.
- Label the taps with either ‘hot’ and ‘cold’ or red and blue colours.
- Using plain coloured plates can help food stand out. Serving drinks in a coloured beaker or mug, rather than a glass, also helps the person see them more easily.

**Stairs and hall**
- Use one colour for all carpets or flooring – patterns can look like an obstacle or step that the person needs to avoid. Shiny flooring can also look wet or slippery, so stick to matt.
- Mark out the edges of any steps or stairs with yellow tape. Paint door frames and light switches a different colour, so they stand out clearly from the wall.

**Living room**
- Good lighting can help a person with dementia find their way around – increase natural light during the day and fit high wattage bulbs in the main lights.
- Remove rugs or stick down the edges – falls are common in the living room. Clear away any wires and remove unnecessary furniture that could easily be tripped over.
- People with dementia can struggle with TV remote controls. Covering up small buttons they don’t use with stickers can really help.
Can you really live well with dementia?

“I know receiving a diagnosis of dementia may be difficult, but your loved one can still live a full, independent life. Following a healthy diet, staying active and keeping up with friends and hobbies will help them live as well as possible, for as long as possible. Focusing on their strengths and what they can still do, rather than on the challenges of dementia, will also help boost their mental wellbeing. Don’t forget our Helpline is always here when you need us.”

Admiral Nurse Vic
Section 3

Your wellbeing

Taking care of yourself often comes way down the list when you’re caring for someone with dementia. But looking after your own wellbeing is just as important. Making time for yourself can improve your mental and physical health, help you feel better equipped to care for your loved one, and improve your relationship with them.

What you need to know

• **acceptance**: you may not think of yourself as a carer. You might feel your role is simply part of your relationship with the person with dementia. But if, for example, you help with their personal care, manage their bills or medication, or stay with them if they can’t be left alone, you are a carer.

• **balance**: to make sure you’re also looking after yourself, you first have to recognise what your needs are. Do you want to exercise? Do you need some time alone every day to read or relax? What about a night out with friends? Once you figure this out, you can ask others for help to rebalance your life.

• **keep checking in**: once a week, for example, do a mini assessment of how you and the person with dementia are coping. “What’s this week been like for us? Are we doing okay? Is it safe for both of us to keep going this way?” If you spot any problems, you can sort them out sooner rather than later.

• **health checks**: if you’re feeling unwell, have any niggling aches or pains, or experience anxiety or depression, don’t ignore it. If you’re unable to look after the person because you’re ill, what might happen? Tell your GP you’re a carer so they can put it on your medical record.

• **extra help**: if you’re finding it tough, it’s important to know what support is available. You could be eligible for funding or respite care (read about the Carer’s Assessment opposite), or you could join local carer support groups to meet others who understand what you’re going through. The dementia specialist Admiral Nurses on our Helpline are here for you too.

I wish I’d known...

“It’s too easy to put your own health on the backburner when you’re a carer. My Admiral Nurse, Amy, was able to pick up on my feelings of guilt at not being able to do enough for my sister, Fiona. She made me realise I needed to take care of my mental health, otherwise I risked being overwhelmed and of no great use to Fiona.”

*Jolyon* helps care for Fiona, who has Alzheimer’s disease.

What helped me most

“Our Admiral Nurse helped me to prioritise my own wellbeing; going to the gym has now become my outlet. It’s so good to get that time to myself and it’s helped enormously with my physical and mental health. I don’t feel like I’m a carer when I’m at the gym – I’m just me.”

*Rachel*, who cares for her mum Brenda.
Quick Q&A

One way to get more support is to have a Carer’s Assessment. Admiral Nurse Kerry takes us through some common questions.

Q. What is a Carer’s Assessment?
A. A Carer’s Assessment looks at how caring affects your life and what you might need to make it easier. It’s often carried out at the same time as a needs assessment (see Section 2 for more information) by a social worker who will ask you about a typical day, what caring responsibilities you have, such as washing or dressing the person with dementia, and whether you get any time to yourself. They’ll also want to know what would happen in a crisis – if you had an accident while out shopping, would anyone know you’re a carer? Who would look after the person with dementia? Try to make some notes about these points before your assessment.

Q. Why have an assessment?
A. After the assessment, the social worker will draw up a care and support plan with ideas to ease some of the challenges you face. You may be eligible to get funding for home adaptations, such as fitting handrails next to the bath, or respite care for your loved one. Apart from funding, this may include details of support groups, help with housework, or tips to look after your wellbeing.

Q. How do I organise a Carer’s Assessment?
A. Contact your GP or local social services. Be aware that your finances will also be assessed to find out if you have to pay partly or in full for any help you need (called ‘means testing’). If you’re not eligible for funding, you should still be given free advice about other ways to find support in your community.

Heard on the Helpline

“I never have enough time for myself.”
Being a carer is physically and mentally demanding, so it’s incredibly important to look after your own health needs. Can you ask someone to sit with the person with dementia while you go out? Make sure you keep any GP or dentist appointments, get regular exercise and keep up with the social activities you enjoy, too. You deserve a life outside of being a carer.

“Who can I ask for help?”
Your own family and close friends may not realise how much you’re doing. Talk to them to see if they can take on some caring tasks. Search online for dementia support groups near you (Facebook is a great place to start), while local libraries or community centres may have more information about support groups. You could also be entitled to respite care – a paid carer who looks after the person at home for a couple of hours every week or longer periods – or your loved one may be eligible for a place at a day centre. Contact your local social services to start the ball rolling.

Life hack

“Having a smartphone is fantastic; you can check your emails, do an online shop, message friends, watch TV. But if you don’t charge it up, it’s basically a coffee coaster. You need to recharge your batteries in the same way – learn what helps to recharge you and make sure you ‘plug in’ to those resources.”

Admiral Nurse Gary
Although your relationship may change, you can still enjoy a close and meaningful connection to the person with dementia. Try these top five tips from Admiral Nurse Vic:

1. **What hobbies did you enjoy doing together?**
   If it was hiking, for example, can you recreate that sense of enjoyment by going for short walks in the park?

2. **You can do other activities to help bring you closer**, such as watching your favourite films, listening to music, or looking at old photos together.

3. **When your partner has dementia**, this can affect your sex life. Some people find their other half wants to be more intimate, while others do not. But you can stay connected in other ways, such as touching or cuddling. As dementia progresses, this may bring up questions of consent. Keep talking to each other or ask a professional for advice to help manage any changes.

4. **If the person gets distressed**, they may start shouting, crying or even behaving aggressively. This may mean they’re trying to communicate something they can’t express. Try to find out why they’re distressed and resolve it, if possible. You may want to practise some calming techniques, such as deep breathing exercises; comfort and reassure them; or you may just need to give them some space.

5. **Try not to let your role as a carer take over** the previous relationship you had with your loved one – always remember the person behind the condition.

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**Heard on the Helpline**

“**I feel guilty all the time – is that normal?**”

Unfortunately, yes. You might feel guilty if you think you’re not living up to what a carer ‘should’ be; if you have to reconsider a promise such as, “I’ll never move you into a care home”; or if you need some time to yourself. But remember you’re only human and you’re doing the best you can. Getting help and support will take some of the pressure off and help reduce feelings of guilt.

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**Watch Chief Admiral Nurse Hilda talking about guilt**
Section 3: Your wellbeing
Managing behaviour changes

Everyone with dementia experiences changes in behaviour, from misinterpreting information to getting agitated around sunset. These changes may be distressing, but there are ways to manage what’s happening.

What you need to know

- **find the cause**: changes in behaviour could be caused by a specific trigger, like a noisy train station or busy supermarket. Ask yourself what happened just before your loved one’s behaviour changed; what were they doing? What was happening around them? Who were they talking to?

- **medical reasons**: agitated or aggressive behaviour is sometimes triggered by feeling unwell or being in pain. If it continues, the person should see their GP.

- **book a check-up**: changes in hearing or eyesight can lead to difficulties with perception and confusion. When did they last have a check-up?

- **extra advice**: if you’re still not sure what’s behind your loved one’s behaviour changes, our Admiral Nurses are here for more help and support.

**“The doctor says my husband has delirium. What is it?”**

Delirium is a state of intense confusion and lack of awareness of their surroundings, often causing distress. It often comes on suddenly and symptoms may include paranoia, restlessness, aggression, confusion about the time or day, and hallucinations. There are many different causes, including infection, dehydration, side effects of some medications or a mini stroke. If your loved one has delirium, they need medical help right away, so contact their doctor (or whoever is responsible for their care). Stay with them and try to keep them calm – distract them by watching TV, offering food or drink, or walking around with them. Delirium can be treated (most people get better within a week) but some people with dementia take much longer to recover.

**“Mum doesn’t recognise me anymore, which is really upsetting.”**

As dementia progresses, many people stop recognising those closest to them. For example, they may think you’re someone else or expect grown-up children to be small again. This could be caused by memory problems or damage to the part of the brain that recognises faces. Try not to correct your mum, as this can be distressing and frustrating for everyone – see the suggestions on page 22 for advice on what to say instead. You could also put up photos that show people getting older or important events in your life together.
Quick Q&A

Admiral Nurse Gary explains a common change in behaviour known as sundowning.

Q. What is sundowning?
A. More than 90% of people with dementia experience sundowning. It means the changes in behaviour that normally happen in the evening, around dusk.

Q. What are the symptoms?
A. People who experience sundowning often become anxious, agitated and confused. They may have a strong sense that they’re in the wrong place and want to, or try to, go home – even if they’re already in their own home.

Q. What causes sundowning?
A. Possible causes are getting tired at the end of the day, feeling hungry or thirsty, darkness falling and the streetlights switching on. These changes can all make the person feel unsettled or as if they’ve forgotten to do something.

Q. How can you manage it?
A. Use distraction techniques, like offering them a drink or going for a walk. Ask them what’s wrong in a low, soothing voice, and listen carefully to their answer. Turn off the TV or radio, as background noise can be overwhelming. Sit with them and hold their hand to comfort them, too.

Q. Can you prevent sundowning?
A. Try to limit naps during the day, so your loved one sleeps better at night. Also limit their intake of caffeinated drinks – switch to decaf coffee or herbal teas. Draw the curtains and put the lights on before it gets dark to soften the shift from day to night-time. Try to create a regular evening routine, such as eating a meal, having a relaxing bath, then dressing for bed.

I wish I’d known...

“Grandad would get very distressed every evening. He insisted he needed to go home, would try to pack a bag and demand to be taken to the airport so he could fly back to India where his parents and siblings, long since passed away, were waiting for him. After calling the Dementia UK Helpline, we learned to acknowledge what Grandad was saying but to quickly change the subject and distract him. We told him it was too late to go to the airport and asked him what he wanted for dinner.”

Liz, whose grandad, Sam, has Alzheimer’s disease
Talking to someone with dementia

You shouldn’t ask people with dementia too many questions, but do you know why? Admiral Nurse Kerry says it’s because a lot of human interaction is based on questions: how are you? What did you do at the weekend? When did Jay come to see you? For someone with dementia, this can be overwhelming and raise their anxiety levels if they don’t know the answers. Try these simple alternatives instead:

**Don’t say**

- “Don’t you know who I am?”
- “Hello Mum – it’s Julie, your daughter.”
- “Did Danny get you those flowers?”
- “Would you like some fruit?”
- “What did you have for lunch?”

**Do say**

- “It looks like you had a sandwich for lunch.”
- “Your grandson Danny came to see you today and left these flowers for you.”
- “I’m going to have fruit for pudding. Would you like some too?”

Want more information on behaviour change?

Watch all of our videos online
“My partner thinks people are stealing from them. How can I reassure them?”

Some common false beliefs and delusions include the person thinking their partner is being unfaithful, that others are stealing from them, that someone wants to harm them, or that they are living in the past. They may be caused by delirium or changes in the brain that happen as part of dementia.

These false beliefs feel very real, so try explaining what’s happening without challenging or correcting them. Say, “I can see you’re upset that someone might have stolen your purse. Let’s see if we can find it in case it was lost accidentally.” You could also agree to keep important things, like keys, in a certain spot.
Dementia can affect family and friends in lots of complex ways, and it will affect everyone differently.

What you need to know

• be open: encourage your family and close friends to talk about your loved one’s dementia. This means you can understand and support each other better

• share: let other family members know what your caring responsibilities involve. Very often, people don’t realise just how much you’re doing

• ask for help: if possible, split your caring tasks among other family members and close friends – it may be useful to create a rota

• avoid taking over: it’s tempting to take charge of all household tasks, but this can make the person with dementia feel a loss of responsibility and independence, and put a strain on your relationship

• be kind to yourself: many people go above and beyond when it comes to looking after a loved one, but never feel like they’re doing enough. Try to acknowledge what you’re doing and how much you’ve achieved

• be prepared: becoming ‘a carer’ doesn’t mean you’re no longer a partner, friend, sibling or child to the person with dementia, but you may need to adapt to a change in your ‘role’. Being prepared can help you understand and accept these changes and maintain positive relationships

Heard on the Helpline

“I feel like I’m doing everything.”

Very often, one member of the family feels as if all the caring responsibilities fall on their shoulders. If other family members can’t (or won’t) help – even after you’ve suggested sharing tasks – you may need extra support. Talk to our Admiral Nurses to find out what’s available near you.

“As a family, we can’t agree on how to look after Dad.”

When family members have different opinions about how their loved one should be cared for, it can be difficult to agree on and plan the next steps. Some may feel excluded from caring or feel uncomfortable volunteering as they don’t know exactly what to do. If you’re struggling to work things out, contact our Helpline or organise a virtual clinic appointment. Our expert Admiral Nurses can help the whole family find ways to care for the person with dementia.
Quick Q&A

If you don’t live near your loved one, it can be difficult trying to care for them from a distance. Chief Admiral Nurse Hilda has the answers to these common questions.

Q. How can I keep the person with dementia safe?
A. The first step is to ask the local social services team to carry out a needs assessment. This will help them work out what equipment or extra support the person needs to live safely and independently at home. See Section 2 for more advice.

Q. What if something happens when I’m not there?
A. Build up a local network of trusted people, such as neighbours, family and friends, who can keep an eye on your loved one and contact you if anything happens. You could give a select few friends a spare key or door code, so they can get in should they need to – just make sure the person with dementia is comfortable with them. It may also help to keep all the information about the person, such as their care needs or doctor’s details, in one place, so it’s easy to find in an emergency.

Q. What happens if/when they get worse?
A. Try to have open discussions with the person with dementia and other family members about future plans, care options, finances and everyone’s expectations to avoid conflict later on. It’s also important to draw up a lasting power of attorney while the person still has capacity – this will allow you to make decisions on their behalf if they’re unable to; see the Legal and financial issues section for more.

Loved, not lost

“If/when a person develops dementia, it doesn’t change the person you love and respect. A grandparent will still be a grandparent, a parent will still be a parent, a friend will still be a friend. People may have dementia but dementia does not have them.”

Rueben, healthcare assistant
5 top tips for talking to children

It can be tough talking about dementia to children, who may be struggling to understand any changes in their parent or grandparent. Admiral Nurse Kerry shares five useful tips:

1. **When you’re explaining what dementia is, be honest about what’s happening and how it may affect your family, but try not to overwhelm them.**

2. **Make time to listen to their concerns.** Give them the chance to open up and reassure them it’s okay to feel a range of emotions.

3. **Create a memory box – put in photos, cards and keepsakes to help them remember their loved one before they were diagnosed with dementia.**

4. **Make new memories, too.** Spend quality time walking, playing board games, or watching some family films together.

5. **Ask their teachers to look out for any behaviour changes, which could be a sign they’re not coping. If you need more support, please call our Helpline.**

**Let’s talk dementia**

We ask children to share their tips for eight to 12 year olds. [Watch now](#)
“One thing we agreed on is that we needed expert help”

“Living with dementia takes up a lot of bandwidth, even when I’m working. I am always worrying about Mum and put a disproportionate amount of hope on the ups, which makes the downs harder to cope with. I’m living with a constant fear of what is next.

“My sister, Shahbanu, and I disagreed hugely about what was happening. I thought she was burying her head in the sand and would get angry when she lost her patience with Mum over her forgetfulness. One thing we agreed on was that we needed expert advice to support Mum as best we could.

“We booked an appointment in Dementia UK’s virtual clinics and spoke to Ruby, an amazing Admiral Nurse. She gave us a framework for how we could best help Mum and ourselves, which made us feel more prepared for what’s to come. Now we know what we’re dealing with, we can all move forward.”

Aqib, who lives with his sister Shahbanu and their mother, Shama, who has Alzheimer’s disease
Section 6

Legal and financial issues

Are you worried about the financial impact of caring for someone with dementia? Perhaps you’re feeling anxious about making future decisions about health, money and long-term care if your loved one can’t do it themselves. Figuring all this out can feel daunting, but we’re here to help guide you through it.

What you need to know

- **help is out there**: many carers don’t know they can claim certain benefits or get financial support, so it really is worth finding out if you’re eligible
- **legal issues**: there are steps you and your loved one should take sooner rather than later, like setting up a lasting power of attorney, to help them feel more confident about the future
- **simplify your finances**: managing money can be tricky — especially if your income suddenly drops — but there are ways to make it easier, such as setting up direct debits for bills
- **ask an expert**: negotiating your way through unfamiliar legal jargon or benefits forms can be overwhelming. Our dementia specialist Admiral Nurses are here for you and have helped hundreds of people make sense of it all

**Heard on the Helpline**

**“Are we eligible for any financial benefits?”**

People with dementia and their carer may be entitled to a significant council tax reduction, Attendance Allowance, disability premium, Universal Credit or Personal Independence Payments. You could also claim a Carer’s Allowance or Carer’s Credit. Receiving benefits will depend on many factors — your loved one will need to meet various criteria, such as state pension age.

**“What is NHS continuing healthcare (CHC) funding?”**

CHC is for adults with long-term complex health needs, including dementia. It’s designed to cover social care costs, such as home carers or care home fees, and is paid for by the NHS. However, the application process is often very difficult, so we strongly advise getting help from a dementia specialist who understands CHC, like our Admiral Nurses.

**Watch Admiral Nurse Kerry talk about CHC**
Quick Q&A

Admiral Nurse Vic answers the most important questions about a lasting power of attorney (LPA).

Q. What is a lasting power of attorney?
A. A person with dementia can choose one or more trusted people to make financial, legal or healthcare decisions for them in the future. This person is called an attorney and is appointed by a formal document called a lasting power of attorney. There are two types of LPA: one for health and welfare, and one for property and affairs.

Q. Why have an LPA?
A. Many people with dementia reach a point where they can no longer make decisions about their own money or healthcare – this is called ‘loss of mental capacity’. Making an LPA before this point means your loved one can make choices about their future care and finances while they’re still able. The attorney is then legally bound to respect their wishes and make decisions in their best interests. But if the person with dementia loses mental capacity and doesn’t have an LPA, it can be very difficult for you to legally make those decisions for them.

Q. How do I set up an LPA?
A. You can contact a solicitor or download and complete the form from the Office of the Public Guardian. Once you’ve filled it in and got it signed by everyone, you need to post the form to the Office of the Public Guardian to be registered.

Q. My dad won’t agree to an LPA – how can we encourage him?
A. Many people, whether they have dementia or not, worry that having an LPA means someone else will make decisions about their health or finances that they don’t agree with. But not having an LPA could mean your loved one doesn’t have any say in what they want – and explaining this could help change their mind. It’s also worth pointing out that an attorney cannot take advantage of the person for their own benefit. If they do, the LPA may be cancelled, and the attorney could be prosecuted.

Heard on the Helpline

“How do I apply for Carer’s Allowance?”
You can claim Carer’s Allowance if you look after someone who receives other benefits, like an Attendance Allowance. But you’re not eligible if you already receive a State Pension and – most importantly – it may affect the benefits your loved one could claim, too. Get help with an application by calling the Carer’s Allowance Unit on 0800 731 0297.

“Quick Q&A”
Admiral Nurse Vic answers the most important questions about a lasting power of attorney (LPA).

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Dementia can change your loved one’s relationship with money. For example, if your partner was diagnosed with dementia while working and needs to leave their job, this could have a huge impact on your income. The condition can change how a person handles money, too; they might start giving it away, for example. Chief Admiral Nurse Hilda shares some tips to help you and the person with dementia manage their finances:

1. **Set up standing orders or direct debits** for all your regular bills. Signing up for online banking also makes it easier to track your transactions.

2. **If you want to limit someone’s spending**, prepaid cards are a great idea. Or you can set a spending limit on debit and credit cards.

3. **Create a household budget** – list all your incomings and outgoings so you know exactly what you’re spending every month.

4. **If you have multiple joint bank accounts**, consider closing some to avoid confusing your loved one.

5. **Separate bank accounts can be helpful** – if the person with dementia has a financial assessment for their own care needs, only their finances will be considered, and not yours.

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**I wish I’d known…**

“The biggest issue I had with CHC was that it was so difficult to find my way around the system. There was a real lack of consistency – if you rang up and spoke to someone you hadn’t been talking to before, they didn’t know who they were dealing with. I wish I’d known how important it was to record every bit of information, otherwise you feel like you are always starting again at the beginning.”

Mary, who cared for her mum who had Alzheimer’s disease
What helped me most

“It was really worth spending the time to get our heads around the benefits available to us as a family after Mum’s diagnosis. Although completing the Attendance Allowance form took time, including a couple of weeks to get all the information together, having it in place then unlocked other benefits for us, such as council tax reductions.”

Claire, who cares for her mum who has dementia

Life hack

“Both you and the person with dementia could be entitled to different benefits that you may not be aware of. Use the government’s online benefits calculators to see what you could claim for.”

Admiral Nurse Gary
Any hospital stay – whether planned or an emergency – can be stressful for the person with dementia, and for you. From packing a ‘grab bag’ to advice on speaking to staff, our tips will help you feel better prepared for a hospital visit.

What you need to know

- **get packing:** it’s a good idea to pack a ‘grab bag’ with all the things the person might need if they’re admitted to hospital unexpectedly. As well as toiletries, include nightwear and supplies of medication. If they’re going into hospital for a planned stay, you might want to pack comforts such as noise-cancelling headphones or a favourite blanket that will help them feel at home

- **essential info:** you could fill in a care passport, which includes all the important information hospital staff might need to look after the person. Go to the next page to download our PDF

- **explain their needs:** check that the hospital knows your loved one has dementia, understands their needs and can communicate with them clearly; and that you’re comfortable that they will care for them properly

- **extra support:** ask if the hospital has a dementia team or an Admiral Nurse who can support both you and your loved one during their stay

- **keep moving:** many people with dementia like to walk around. If they’re prevented from doing so, they may become distressed. Talk to hospital staff to see if it’s okay for your loved one to walk around the ward, day room, or to go for a coffee in the hospital grounds with a visitor. And ask if they have any resources to help reduce your loved one’s anxiety or distress

"How can I help Mum feel more comfortable in hospital?"

Involve the person with dementia in packing their bag by discussing what they’d like to take, such as clothes, books, magazines or a favourite shower gel. Label all their items clearly so they don’t worry about losing them, and advise them to leave valuables (like wedding rings) at home. If you can’t visit them, ask the ward staff if there is a good time to phone or video call them. Some hospitals also take part in John’s Campaign, which allows friends and family of people with dementia to visit at any time. Agree with your family and friends who will visit and when, so your loved one is never alone for too long.

"Do I need a legal power of attorney (LPA) if they go into hospital?"

Not necessarily. If the person with dementia is still able to make their own decisions (ie if they have mental capacity) then an LPA isn’t needed. But if their condition progresses, or their mental capacity comes and goes, then a trusted family member or friend may need to make decisions about their healthcare. It’s a good idea to make an LPA soon after diagnosis to avoid problems further down the line.
Quick Q&A

Most people receive excellent care in hospital, but if you feel your loved one isn’t being properly cared for, Admiral Nurse Kerry recommends these steps to make a complaint.

Q. Have you spoken to their named nurse?
A. This is usually the first step to resolving any issues. Check that they know about the person’s dementia and are following their care passport, if they have one. If this doesn’t work, ask to speak to the ward manager.

Q. What’s your specific complaint?
A. Write down your concerns, including dates, days, times, what happened and who you spoke to. Then you can be precise when making a complaint. For example, ‘Yesterday, Mum’s breakfast was cleared away before she had finished eating. I spoke to staff at 11am who did not know she had dementia and needed more time. But then her lunch was also cleared away early.’ Be clear about what you want them to do to fix the situation.

Q. Has this solved the problem?
A. If you still have concerns, or the problem keeps happening, speak to the hospital’s Patient Advice and Liaison Service (PALS). They support patients and families with resolving issues during or after a hospital stay and can advise on making a formal complaint.

Create a care passport
Download our PDF tailored for families living with dementia.

Life hack

“Set up a family WhatsApp or group chat so everyone gets the same updates about the person with dementia at the same time. The main carers don’t have to be in the group themselves – to save them from too many notifications buzzing on their phone all day – and it can be really handy for delegating responsibilities too.”

Admiral Nurse Gary

What helped me most

“Find out if there’s a dementia specialist in the hospital, either before the person goes in or as soon as they’re admitted. Admiral Nurse Kerry came into our lives a few hours after Dad was taken to A&E. She gave us practical help, regular updates, messages of reassurance and advice, and support in finding him a care home place. But crucially, she made us feel like someone cared about our dad, not just that he was being cared for.”

Neil, whose father had Alzheimer’s disease
When people with dementia stay in hospital, they may lose their appetite. Admiral Nurse Vic has these top tips to keep them eating and drinking well:

1. **Tell nurses and other care staff** about any allergies or food and drink they like and don’t like.

2. **Ask if there are picture menus** so the person with dementia can easily see what food is available, especially if they struggle with words.

3. **Is there a snack menu?** They may find it easier to eat smaller, more regular meals rather than three large meals a day.

4. **Some hospitals have a ‘red tray scheme’** which shows the person may need more help during mealtimes. Find out if you can be there to help your loved one with eating meals.

5. **If the person has trouble chewing or swallowing,** ask if they can have softer food or thickened drinks to make it easier for them to eat. The hospital will be able to refer them to a speech and language therapist (SALT) or dietitian assessment to help with this.

**I wish I’d known…**

“Check if doctors or nurses have changed your loved one’s medication, like adding something new, or increasing or lowering a dose. This can have a huge impact. I remember Nana suddenly changed: she was more sleepy and hardly spoke. My family had no idea this was because her meds were different. You need to ask questions: why have they been changed? What are the benefits? Is it just while they’re in hospital? You all deserve to know what’s happening.”

*Sarah, whose grandmother had Alzheimer’s disease*
“Why do we need an assessment before my partner is discharged?”

This is part of the discharge planning process to make sure your loved one has all the support they need when they leave. A discharge plan looks at a person’s needs, where they will live, any equipment or home adjustments they need and who will look after them. It might also consider whether your family needs more support to care for them. The plan is often reviewed after discharge to make any changes.

If your loved one is leaving hospital at the end of their life, turn the page for our Understanding your care options section.
Section 8

Understanding your care options

It’s not easy looking after someone living with dementia and at some point, you may need extra help. That could be hiring carers to support your loved one at home or moving them into a care home. You may also be looking at end of life care. So, what’s available and how do you make those decisions?

What you need to know

- **it’s personal:** there’s no ‘one size fits all’ when it comes to dementia care. Every family needs different levels of support at different times, so it’s important to get advice, and realistically work out what’s best for your loved one and for you
- **cost:** paying for care can be a huge concern. Find out if you’re eligible for any funding by asking social services for a needs assessment. You could also apply for a Carer’s Assessment; see the Your wellbeing section for more advice
- **quality care:** every care home in England is inspected by the Care Quality Commission (CQC). You can see all the ratings on their website. Care agencies that provide carers are also regulated by the CQC. Details of care inspectorates in Northern Ireland, Scotland and Wales are on the back page
- **end of life needs:** recognising when someone with dementia is nearing the end of their life can help you plan appropriately, helping to ensure their care needs are met, if possible, and final wishes are carried out respectfully. Read our leaflet for more information

So, what are your options?

For more information on dementia care at home, in a care or residential home, or in hospital, head to our website. And visit Care Home for trustworthy reviews and advice.

Heard on the Helpline

“Is there a right time to move my loved one into a care home?”

Some people set themselves a personal limit, such as ‘When Mum becomes doubly incontinent, she’ll be better cared for in a home’. But that point varies for everyone. It could be following discharge from hospital after an emergency when they need specialist support, or if the person’s dementia progresses to the stage where they need 24-hour care. A home may also be more practical if you, or their regular carer, are unable to keep looking after them.

Life hack

“When you book viewings at care homes take note of other things apart from the décor and furniture – are the residents dressed appropriately? Are their faces and nails clean? Are they engaged in some sort of meaningful activity? Listen to how the staff talk to the residents – is it respectful? And do they look happy in their job? All this can help you make the right decision.”

Alison, who works in a dementia care home
Quick Q&A

If you’re thinking about end of life care, or your loved one is close to dying, Chief Admiral Nurse Hilda says these are the key questions to ask.

Q. Has the person with dementia got an Advance Care Plan?
A. An Advance Care Plan (ACP) sets out how your loved one wants to be cared for as they approach the end of life. It often includes their wishes for medical treatment, plus details like funeral plans or caring for pets. If possible, it’s important to make an ACP while the person can still make decisions, so you know you’re following their wishes.

Q. Does the plan cover where they’d like to die?
A. If you’ve not yet discussed where the person would like to die, have that conversation sooner rather than later. Some people prefer the idea of dying in their own home; others may need to be in hospital to meet their needs at this important time. They could prefer the idea of a care home, or the care of a hospice. Whichever feels right for your loved one, make sure their requests are in the ACP – and followed as closely as possible.

Q. What about advanced decisions, or a ‘living will’?
A. While an ACP is not a legal document, an advance decision to refuse treatment (ADRT) – also known as a living will – is legally binding. This sets out what medical treatment your loved one does not want, such as ventilation or resuscitation. It needs to be signed by the person and a witness.

Q. Have you got support during this time?
A. Dementia is a progressive condition, so everyone with the diagnosis will die with it or from it. If your loved one is nearing the end of their life, you will need plenty of support. Keep talking to family, friends or a professional counsellor, or call our Helpline. Our Admiral Nurses can offer support and put you in touch with grief charities or local bereavement support groups.

Heard on the Helpline

“How can we choose the best care home?”

The first step is to involve the person with dementia as far as possible. Do you know what type of home can best meet their needs now, and what support they might need in the future? Ask friends or local dementia support groups for any recommendations. You can also ask social services or the Elderly Accommodation Counsel for a list of care homes. Finally, arrange viewings for you and your loved one. Talk to other residents and their families to get a feel for whether the home suits the person’s needs.

“I feel like a failure for moving my loved one into a home.”

Accepting that you need more help looking after someone with dementia is a big step. It doesn’t mean you’ve failed or given up on them. In fact, finding the right home for your loved one can provide them with vital care and social contact, and give you back some time to resume the activities you enjoyed before becoming a carer. It will feel like a major transition, and some people with dementia may be reluctant to move at first, but it can have huge benefits for everyone involved. If you’re struggling with the change, our Helpline nurses are here to offer support.
At-home carers or a care home?

There are so many care options to choose from that it can be hard to know what’s best. We’ve broken down the key differences between at-home support and care homes to help make your decision a little easier.

<table>
<thead>
<tr>
<th>Will it…?</th>
<th>At-home carers</th>
<th>Care home</th>
</tr>
</thead>
<tbody>
<tr>
<td>... relieve some of the strain on you?</td>
<td>Professional carers can help with challenging tasks such as lifting, dressing or personal care.</td>
<td>Moving your loved one into a care home can relieve you of some of the day-to-day stresses of caring, but if it’s far away, expensive, or a poor fit for your loved one, that could be a strain on you.</td>
</tr>
<tr>
<td>... keep them in their own home for longer?</td>
<td>This choice means your loved one will stay at home – as their dementia progresses, just keep checking their needs are being met, they’re still able to live safely at home, and compare the costs to other options.</td>
<td>As dementia progresses, the person may need specialist, round-the-clock care that isn’t possible in their own home.</td>
</tr>
<tr>
<td>... keep them connected to their community?</td>
<td>If your loved one stays at home, they can keep up with the activities they enjoyed before and see local friends and family.</td>
<td>If your loved one was becoming isolated at home, a care home can provide a new community that understands their needs.</td>
</tr>
<tr>
<td>... provide 24-hour care for your loved one?</td>
<td>Some home carers provide live-in care, or you may be able to use an agency that provides carers to work shifts, including nights.</td>
<td>If your loved one is in a home, they will be looked after both day and night.</td>
</tr>
<tr>
<td>... give them specialist dementia care?</td>
<td>Make sure any carers are trained in dementia care and find out what caring tasks they are able to do.</td>
<td>Ask how many dementia specialists work in the home, about their training (what they’ve done and how regularly), and how much contact they’ll have with your loved one.</td>
</tr>
<tr>
<td>... support someone with young onset dementia?</td>
<td>You can ask for carers who specialise in young onset dementia, but they are quite rare.</td>
<td>Not many homes care for people with young onset dementia, so finding age-appropriate care can be difficult.</td>
</tr>
<tr>
<td>... strengthen your relationship?</td>
<td>Being relieved of some of your caring duties can make the time you have together more enjoyable.</td>
<td>If you’re no longer responsible for tiring or stressful tasks, you can focus on your loved one during visits.</td>
</tr>
<tr>
<td>... improve everyone’s quality of life?</td>
<td>Giving you some much-needed time to yourself, or reducing isolation if they live alone, can help everyone live better with dementia.</td>
<td>Once you’ve adapted to the change, a care home can lead to better health and quality of life, with more time for meaningful activities, for everyone.</td>
</tr>
</tbody>
</table>
“Go with your gut – and don’t settle”

“I wanted Mum to live in her own home for as long as possible, so I tried to make that happen. As her dementia progressed, I was lucky enough to be able to hire 24-hour carers for her. But I still felt like I was always on call, as the carers would constantly ring me. Eventually, I accepted that it was time for Mum to go into a home. While I was trying to find one, I found it best to go with my gut.

“My advice is to look at as many homes as you can and don’t stop until you find the right one. I worried I was being too picky, but I knew what I wanted – and I wasn’t prepared to settle. Now I’ve found a home that suits my mum perfectly and I’m so pleased I didn’t give up.”

Penny, whose mum, Rosemary, has mixed dementia. The pair now host the ‘Discovering Dementia’ podcast
Notes

As you go through this guide, you may want to make notes or remember an important phone number – you can use this section to jot down anything below.
I live with dementia

Find out more about our campaign for everyone affected by dementia.

Visit now
Contact us

To speak to a specialist dementia nurse, call our Dementia UK Helpline free on 0800 888 6678 (Mon-Fri 9am-9pm, Sat-Sun and bank holidays 9am-5pm – closed 25th December) or email helpline@dementiauk.org

Sometimes our Helpline can get busy. If you cannot get through straight away, there is an option to leave a voicemail and an Admiral Nurse will get back to you within 48 hours.

To book a free video or phone appointment in our virtual clinics at a time and date that suits you, visit dementiauk.org/book-a-clinic-appointment

Our free online advice and information is written by our Admiral Nurses and can help you with everything from getting a diagnosis to legal and financial support.

You can find all of our resources at dementiauk.org/get-support

Some of our information leaflets are also available in other languages, including:
Bengali
Chinese (traditional and simplified)
Polish
Punjabi
Welsh

“When I called the Helpline, I broke down. I shared all my worries and for the first time, I felt listened to and understood.”

Denise, whose mum had Alzheimer’s disease
More useful contacts

Here are some of the most important contact details from each section. Cut this page out and pop it on your fridge so you have them to hand for any emergencies. And if you’ve got any questions, you can always call the Helpline or book an appointments in our virtual clinics. Our dementia specialist Admiral Nurses are happy to help.

Organisations that support carers

**Carers UK**
- Phone: 0808 808 777 (Mon-Fri 9am-6pm)
- Website: [carersuk.org](https://carersuk.org)

**Carers Trust**
- Phone: 0300 772 9600 (Mon-Fri 9am-5pm)
- Website: [carers.org](https://carers.org)

**Deep**
- Website: [dementiavoices.org.uk](http://dementiavoices.org.uk)

Respite care and short breaks

**Care Home**
- Website: [carehome.co.uk](https://carehome.co.uk)

**Revitalise**
- Website: [revitalise.org.uk](https://revitalise.org.uk)

Living aids and home adaptations

**Living Made Easy by the Disabled Living Foundation**
- Phone: 0300 999 0004 (Mon-Thurs 10am-4pm)
- Website: [livingmadeeasy.org.uk](https://livingmadeeasy.org.uk)

**Healthcare Pro**
- Website: [healthcarepro.co.uk](https://healthcarepro.co.uk)

**AlzProducts**
- Website: [alzproducts.co.uk](https://alzproducts.co.uk)

Young onset dementia

**Young Dementia Network**
- Website: [youngdementianetwork.org](https://youngdementianetwork.org)
- Email: youngdementianetwork@dementiauk.org

Legal and finance

**The government benefits calculator**
- Website: [gov.uk/benefits-calculators](https://gov.uk/benefits-calculators)

**Carer’s Allowance Unit**
- Phone: 0800 731 0297 (Mon-Fri 8am-6pm)
- Website: [gov.uk/carers-allowance/how-to-claim](https://gov.uk/carers-allowance/how-to-claim)

Support and protection at work

**Acas**
- Website: [acas.org.uk](https://acas.org.uk)

**Access to Work grant**
- Website: [gov.uk/access-to-work](https://gov.uk/access-to-work)
  (England, Wales and Scotland)
- Website: [nidirect.gov.uk/articles/access-work-practical-help-work](https://nidirect.gov.uk/articles/access-work-practical-help-work) (Northern Ireland)

Care home inspections

**Care Quality Commission** (England): [cqc.org.uk](https://cqc.org.uk)

**Care Inspectorate** (Wales): [careinspectorate.wales](https://careinspectorate.wales)

**Regulation and Quality Improvement Authority** (Northern Ireland): [rqia.org.uk](https://rqia.org.uk)

**Care Inspectorate** (Scotland): [careinspectorate.com](https://careinspectorate.com)

All information correct as of Jan 2023
Thank you for reading this guide

Bookmark this guide in your browser, save it to your desktop or smartphone files or print a copy to have handy.

If you have any questions or need further advice, please speak to one of our specialist dementia nurses. We’re here for you:

📞 Call our free Helpline
0800 888 6678

✉️ Email us at
helpline@dementiaku.org

📅 Book a virtual appointment
dementiaku.org/book-a-clinic-appointment

If you have found the information in this guide useful and would like more families to benefit from our advice and support, please consider making a kind gift.

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Thank you.