Young onset dementia pathway

Diagnosis and support for people with young onset dementia and their families

Recommendations from the Young Dementia Network Steering Group
Summary

This document brings together the experience and expertise of people affected by young onset dementia and the people who work with and support them. It highlights that the impact of dementia can be very different depending on your stage of life and that because of this, services must be tailored to meet those needs. Currently, this is rarely the case, and this means that a younger person can take twice as long to be diagnosed than an older person. Once they receive this diagnosis they are often left with no appropriate support and no one with the specialist knowledge to guide them through the journey ahead.

From first contact with a GP to end of life care, that journey can be very different for a younger person. They are likely to be in work, have financial responsibilities and dependent children. They may have a rarer form of dementia which means they are part of an even smaller minority. A dementia diagnosis is always devastating. But for a younger person, that diagnosis means that life and future plans change beyond recognition.

Together we have identified a series of recommendations which are designed to meet those different needs. These recommendations will guide policy makers, commissioners and practitioners in designing and delivering services. Most importantly, we hope they will empower younger people with dementia and their families to demand the services to meet their needs.
Recommendations

Recommendations for policy makers and commissioners

- Young onset dementia services do not need to cost more – they can be formed through drawing together a team from existing dementia services.

- Commissioned services must be designed to address the specific needs of younger people with dementia and their families. This means that providers must be able to demonstrate how their service is adapted for younger people. In particular, offering or signposting to age-appropriate activities and support to maintain physical, cognitive and mental health and social engagement for people at all stages of young onset dementia; and specialist support for families.

- Services must be commissioned to meet demand. The number of people living with young onset dementia in each geographical area can be calculated using evidence-based prevalence figures and dementia databases. This is best done by understanding the age at diagnosis of all people living with dementia in order not to miss those who are now over 65.

- Dedicate commissioning to developing and supporting existing services to allow sustained clinical experience and development of continuity and co-ordination across different service providers. Ongoing care needs to be specialist or shared, not from GP alone.

- Appoint a dementia lead in the Integrated Care Board to link with a multi-agency partnership responsible for identifying needs and improving services for younger people with dementia.

- Adopt a specific strategy for young onset dementia and implement a joined-up care pathway across primary care and specialist services, including social care.

Recommendations in detail

- Positive contact with my GP which reduces the time taken to receive the correct diagnosis
  - GPs to have an awareness of young onset dementia and the variety of symptoms with which it can present including atypical ones. This could improve recognition by GPs of key symptoms that may indicate a referral to specialist services is required
  - clear referral routes from the GP to specialist diagnostic services

- A clear and collaborative diagnostic process which is young onset specific
  - ensure a defined local care pathway for younger people with possible cognitive impairment with special attention to transitions between primary care and assessment services; and assessment services and post-diagnostic support services
  - provide access to a specialist with expertise in the diagnosis of young onset and rare dementia, to improve standards of assessment and diagnostic accuracy

- Specialist support and information which meets our changing needs
  - A specialist dementia practitioner who has knowledge, training and experience of young onset dementia and its impact on the person with the diagnosis and their family, as well as knowledge of local and national service provision
Living with young onset dementia

- for the specialist dementia practitioner, clinicians, social services and charitable organisations to work closely together to provide a comprehensive service for the person with young onset and their families
- access to age-appropriate and meaningful activities which enable younger people and their families to remain engaged in their communities, retain and develop relationships and social connectedness. Age-appropriate means that the content, location and timings of activities are designed to be beneficial and accessible for younger people
- advice which enables the person to plan for common issues such as: how to manage or leave employment, how to manage financially, future planning and for the end of life
- access to age-appropriate and local short-term and long-term care, based in their own home or in residential care depending on the person’s wishes and needs

Recommendations by role

We recommend GPs should

- be aware of referral routes to services
- be aware of the young onset dementia lead for their area
- when the person has received a diagnosis, develop, agree and regularly review a shared care plan which ensures that they are linked into local support services. This is especially important when a diagnosis was given in a regional or national centre

We recommend clinicians based in diagnostic services should

- have a named young onset dementia lead
- have knowledge of the varied impact of a diagnosis of young onset dementia on the person and their family
- have access to the full range of assessment, investigations and personnel necessary to make a diagnosis and provide ongoing post diagnostic support

We recommend the specialist dementia practitioner role should

- have specialist knowledge, skills and experience of the impact on young onset dementia. This will include a knowledge and understanding of rarer forms of dementia, as well as the impact of a diagnosis at a younger age for the person diagnosed and their family
- provide information, practical and emotional support for family members, as well as connect them to others for mutual support
- provide continuity of support, and enable the person living with young onset dementia to understand their condition and actively engage in their care plan and journey
- act as a coordinator of services, organisations and people, helping to connect the person living with young onset to local groups networks and support services. They will also act as a liaison between clinical services and the person’s community and home
- may be delivered through a variety of organisations depending on local infrastructure and funding arrangements. For example (but not limited to) primary care, voluntary sector or be based within local mental health teams or neurology services

We have used the term ‘specialist dementia practitioner’ throughout this pathway but people in this role may have a different / alternative job title. Titles such as case manager or key worker are also commonly used.
## A young onset dementia pathway

<table>
<thead>
<tr>
<th>Positive contact with my GP which reduces the time taken to receive the correct diagnosis</th>
<th>Services to support these needs</th>
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<tbody>
<tr>
<td><strong>Person with young onset dementia and their family carers</strong></td>
<td>GP aware of young onset dementia and appropriate local referral routes</td>
</tr>
<tr>
<td>We have the right to an early and accurate diagnosis</td>
<td>• GP knowledgeable about young onset dementia and the variety of symptoms which can present, which may differ from those with later onset dementia</td>
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<tr>
<td>My GP listens to me and doesn’t discount dementia because of my age</td>
<td>• GP has a low threshold for referral if there are ‘red flags’</td>
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<tr>
<td>I can talk to my GP about how young onset dementia could affect me, or my family</td>
<td>• GP listens to the person and family members to make thorough assessment of symptoms including baseline investigations, physical examination, assessment of activities of daily living and a brief cognitive assessment</td>
</tr>
<tr>
<td>My GP knows about the referral process for young onset dementia, and it is explained to me. I’m able to record these conversations so I can refer to them later</td>
<td>• GP excludes a potentially treatable illness or reversible cause of the ‘dementia’</td>
</tr>
<tr>
<td>I’m told about the Young Dementia Network, and local young onset support services who can support me</td>
<td>Referral is made to the locally agreed young onset referral care pathway</td>
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<tr>
<td>I can participate in counselling tailored to my age and situation, in preparation for the journey ahead</td>
<td>Access to pre-assessment counselling</td>
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<td></td>
<td>• pre-diagnostic review/support and counselling</td>
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<td></td>
<td>Also see ‘Diagnosing Well’ - NHS Well Pathway</td>
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<td></td>
<td>Information is provided, including</td>
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<td>• reason for referral</td>
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<td>• expectation/possible outcome/consent</td>
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<td></td>
<td>• implications for other areas of life</td>
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### A clear collaborative and specialist diagnostic process

<table>
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<tr>
<th>Person with young onset dementia and their family carers</th>
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<tr>
<td>We have the right to an early and accurate diagnosis</td>
<td><strong>Making the diagnosis in specialist services</strong></td>
</tr>
<tr>
<td>We have the right to receive evidence based, appropriate, compassionate, and properly funded care and treatment, from trained people who understand us and how dementia affects us. This must meet our needs, wherever we live</td>
<td>Clinicians based in memory clinic or specialist services / departments where diagnosis takes place are knowledgeable and skilled regarding young onset dementia and the impact the diagnosis can have on the person diagnosed and their family</td>
</tr>
<tr>
<td>I know who my young onset dementia lead is and am kept updated about the diagnostic process</td>
<td>There are designated young onset dementia leads in each memory assessment service / diagnostic department</td>
</tr>
<tr>
<td>I’m offered the same support as older people who are going through the process of being diagnosed with dementia. For example, I might need a family member or close friend to attend appointments with me</td>
<td><strong>Comprehensive assessment is performed by a commissioned multi-disciplinary team</strong> of key professionals including neurology, psychiatry, nursing, clinical psychology and neuroradiology including access to allied services as appropriate eg occupational therapy, speech and language therapy, social worker, physiotherapy and nutritionists and dieticians as necessary</td>
</tr>
<tr>
<td>If I’m supporting someone through diagnosis, I can speak to someone who understands my concerns and can answer questions I may have about what a dementia diagnosis at this age might mean for my family / friend</td>
<td>The team should have the opportunities for multi-disciplinary case discussion</td>
</tr>
<tr>
<td><em>Pre-assessment counselling is routinely provided to establish informed consent, find out what someone wants to know about their diagnosis, who else they would like to be present when it is given, and if there is someone else, they would like to be informed</em></td>
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**Key elements of diagnostic assessment**

- thorough history taking
- speaking to someone who knows the person well
- neurological examination
- blood screen for rare cause
- neuroimaging
- advanced cognitive assessment / neuropsychology
- consider EEG, CSF analysis, genetic testing and other biomarkers
### A clear collaborative and specialist diagnostic process continued

<table>
<thead>
<tr>
<th>Services to support these needs</th>
<th>Person with young onset dementia and their family carers</th>
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</table>

#### Feeding back diagnosis

**Diagnosis is given in a confidential setting and is delivered sensitively.**

There should be a recognition that the person diagnosed with young onset dementia and their family may still be in work, have dependent young children or other care responsibilities, so may have different questions and responses to the diagnosis.

Enough time should be allocated in order to answer these questions.

Depending on the persons’ wishes, they will receive full feedback of the results of their assessment and an explanation about how the diagnosis was reached as a result and should be provided with prompt follow-up.

Where a diagnosis is provided in a national or regional centre, there must be a clear ‘exit plan’ which links the person back into their local services.

#### Introduction to specialist dementia practitioner

- Specialist dementia practitioner works in collaboration with the diagnostic team and introductions can be made at the time of diagnosis or when appropriate for individual.

- Clear information is provided about next steps and how they can be contacted.

- Working together with clinicians to ensure a joined-up service.

Also see ‘Supporting Well’ - NHS Well Pathway.
<table>
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<tr>
<th>Specialist support and information which meets our changing needs</th>
<th>Services to support these needs</th>
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<tbody>
<tr>
<td><strong>Person with young onset dementia and their family carers</strong></td>
<td><strong>A regular review with the person, their family, the clinical team and others involved in their care is established</strong></td>
</tr>
<tr>
<td>We have the right to be respected, and recognised as partners in care, provided with education, support, services, and training which enables us to plan and make decisions about the future</td>
<td>The individual and family should have access to:</td>
</tr>
<tr>
<td>I’m contacted by a specialist dementia practitioner who, if I choose, meets with me to discuss what dementia at my age will mean for me and my family. I have the opportunity to ask about the impact on my job, mortgage and/or other financial commitments, childcare and any other concerns I might have</td>
<td>● information about the diagnosis</td>
</tr>
<tr>
<td>I have the opportunity to learn more about my type of dementia, which because of my age, is more likely to be a rarer form</td>
<td>● evidence based treatment and support eg pharmacological</td>
</tr>
<tr>
<td>If I’m supporting someone with dementia, the specialist dementia practitioner also supports me</td>
<td>● eg Cholinesterase inhibitors and / or memantine, adapted cognitive stimulation therapy, cognitive rehabilitation and counselling, relationship or family-centred work</td>
</tr>
<tr>
<td>If I / we have children or young people living at home, the specialist dementia practitioner discussed family needs and relationships and can direct us to any support we need</td>
<td>● information and advice about age-appropriate social, physical and creative activities and peer support</td>
</tr>
<tr>
<td>Also see ‘Living Well’ - NHS Well Pathway</td>
<td>● allied health professionals as indicated, including clinical psychologists, speech and language therapists and occupational therapists, who can offer specialised approaches</td>
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<td></td>
<td>● Admiral Nurses or specialist dementia nurses who can work intensively with families who are facing a difficulty managing the effects of the diagnosis</td>
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<td></td>
<td><strong>Specialist dementia practitioner engages with individual and their family/friends to understand their needs</strong></td>
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<tr>
<td></td>
<td>● discuss life, achievements, abilities, relationships and aspirations pre and post-diagnosis</td>
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<tr>
<td></td>
<td>● discuss preferred / usual ways of coping and their potential support needs</td>
</tr>
<tr>
<td></td>
<td>● gain understanding of employment status, financial situation, care responsibilities and family circumstances</td>
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<tr>
<td></td>
<td>● meet family members and support circle to understand their perspective and circumstances</td>
</tr>
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</table>
Living with young onset dementia

Person with young onset dementia and their family carers

We have the right to be recognised as who we are, to make choices about our lives including taking risks, and to contribute to society. Our diagnosis should not define us, nor should we be ashamed of it.

I work with my specialist dementia practitioner to prepare for what lies ahead. I am supported to plan my life and what I want for my future including the end of my life.

They provide information about practical issues like financial and care planning, as well as helping me to talk about my condition with my children, family and employers.

I know I can contact my specialist dementia practitioner and they can point me in the direction of other helpful services.

I want to continue my social life, relationships and activities and hobbies. My specialist dementia practitioner helps me to adapt so I can keep doing the activities that I am interested in.

I’m introduced to new activities and social contacts/groups that are more specific to dementia when the time is right for me.

Services to support these needs

Regular contact with joint reviews with a core team including a specialist dementia practitioner and Clinical team with access to specialist nurses, occupational therapy, clinical psychology, speech and language therapists and social work support as necessary.

Review response to medication

Assess needs of the younger person and their family and support family members.

Signpost the younger person to age-appropriate meaningful activities including those that enable the person to maintain identity and continue to make a contribution to society.

Specialist dementia practitioner tailors a plan with the individual and their family which is appropriate for their stage in life.

- outline needs, goals and aspirations and routes to achieving these
- facilitate access to community services and opportunities, both dementia specific and otherwise
- provide or facilitate access to information about young onset dementia and rare forms of dementia and how to communicate the diagnosis to children, family, employers and friends
- provide or facilitate access to specialist advice and support about young onset specific needs eg employment, mortgage and financial obligations and future financial planning
- provides support for advanced care planning
- provide advice and emotional support to family members and others closely involved in care
- link to health and social services to ensure plans are aligned

Also see ‘Living Well’ – NHS Well Pathway
### Towards the end of life

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<tr>
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<tr>
<td>We have the right to be respected, and recognised as partners in care, provided with education, support, services, and training which enables us to plan and make decisions about the future</td>
<td>Regular reviews are adopted as the person reaches the end of their life and support is tailored to provide</td>
</tr>
<tr>
<td>I’m supported to think about and prepare for the end of my life, including talking about what I do and don’t want</td>
<td>- access to short term care that gives family members a break from caring (including day and residential care)</td>
</tr>
<tr>
<td>The support I get is tailored to me, depending on if I live alone or with my family</td>
<td>- acute care in dementia friendly wards which comply with John’s Campaign so a family member can stay with the person during their stay</td>
</tr>
<tr>
<td>The family and friends of the person with dementia are given information and support to help them to prepare for the end of life</td>
<td>- age-appropriate long-term residential care within a reasonable distance from home</td>
</tr>
<tr>
<td>Also see ‘Dying Well’ - NHS Well Pathway</td>
<td>- support to stay in the home if this is what the person wants</td>
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<tr>
<td></td>
<td>- age-appropriate palliative and end of life care</td>
</tr>
<tr>
<td></td>
<td>- bereavement support for families and carers</td>
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</table>
Evidence for our recommendations

Current research is improving but remains behind best practice. The evidence base regarding needs of younger people with dementia and the needs of their families is better developed, although gaps remain. There remain significant gaps in understanding current prevalence rates for young onset dementia in the UK, the evaluation of services and interventions and their impact and cost effectiveness. Where there is recent research evidence, we have included it. Our recommendations in some cases are based upon what people affected by dementia tell us works as well as professional expertise.

Positive contact with my GP which reduces the time taken to receive the correct diagnosis

Dementia teams have become ‘ageless services’ in which staff may not often see people with young onset dementia and may feel unskilled. Many younger people with cognitive impairment are referred as a result of local policy with the Integrated Care Board to adult mental health services who lack expertise in diagnosis of dementia (Rodda and Carter, 2015).

UK evidence and European evidence points to delays in diagnosis for young onset dementia compared to later onset dementia ie four years compared to two years (van Vliet et al, 2013) and lack of a clearly defined local referral pathway is one of the primary causes of delay in diagnosis (Harris et al, 2004).


Living with Young Onset Dementia and actively shaping dementia research through PPI - The Angela Project

Keith Oliver, Dr Mary O’Malley, Prof Jacqueline Parkes, Dr Vasileios Stamou, Dr Jenny LaFontaine, Prof Jan Oyebode & Dr Janet Carter. (Dementia, 2020, 19 (1): 41-48) doi.org/10.1177/1471301219876414
A clear, collaborative and specialist diagnostic process

Young onset dementia has a vast differential diagnosis. It includes not only atypical dementias but also rare sporadic and hereditary diseases. There is clear evidence that younger people are frequently misdiagnosed with functional and other psychiatric conditions and/or are under investigated for rare cause dementia.


Memory Services National Accreditation Programme (MSNAP) (2020) Standards for Memory Services (7th Edition)

Feeding back the diagnosis

Health professionals can be reluctant to speak openly and honestly with patients and their families about dementia, with some refraining from using the ‘D’ word. Although initially discussing the diagnosis may be distressing, evidence suggests most people prefer to know if they have dementia in order to access appropriate support and treatment and to plan for the future. This matter of choice is established at the pre assessment counselling stage.


DoH (2009) ’National Dementia Strategy’; see also Dementia Challenge 2020

Benefits of a timely diagnosis. Dementia Action Alliance report for primary care

Specialist support and information which meets our changing needs

The support needs of younger people with dementia are different from those of people who are older because their diagnosis is ‘off time’, their symptoms are more likely to reflect rarer dementias and their life stage differs. They are more likely to be physically active, have fewer co morbidities, be employed, have responsibilities to children / young people or elderly parents, have more financial outgoings, and have a greater loss of future. Related needs are hard to meet through ‘all age’ dementia services. Only about one in five younger people with dementia in the UK receive specialist young onset dementia services yet the quality indicators suggest people with the condition are best supported by specialist teams.

What are the experiences of people with dementia in employment? Chaplin R & Davidson I. (2016). Dementia, 15(2), 147 – 161


**Cognitive Stimulation Therapy**


British Journal of Psychiatry, 183: 248 - 254

**Evidence base for age-appropriate meaningful social activities**

Football and dementia: A qualitative investigation of a community-based sports group for men with early onset dementia. Carone et al. (2014)

Dementia. E-Pub


**Additional references**

Dementia Action Alliance (2017) - We Statements

This pathway was produced by the Young Dementia Network steering group. The Network is hosted by Dementia UK and is a collaboration between people affected by and working in the field of young onset dementia. It campaigns to bring about improvements for people living with young onset dementia now and in the future.

You may find the following Young Dementia Network resources useful:

- **Decision-making guide for GPs** - a guide created by the Young Dementia Network and endorsed by the Royal College of GPs to aid the recognition of the symptoms of dementia in younger people

- **Diagnosis experience checklists** - for healthcare professionals who are involved in the process of assessing and diagnosing dementia to help them to better understand the needs of younger people going through this process. There is also a checklist for use by people who may have young onset dementia and their family members

- **Eight key needs of people with young onset dementia and family members** – videos created as part of The Angela Project

- **Supporting implementation of the NICE guideline for young onset dementia** - supports implementation of the NICE dementia guideline and sets out the tailored approach needed at key stages of diagnosis and support

- **Young onset dementia ID cards** – cards that can be used when out and about when people may need help and support created by the Young Dementia Network

**Web:** youngdementianetwork.org  
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