1. Purpose of the report

To provide the Health and Wellbeing Board with an update on two reports recently published by Healthwatch Derbyshire (HWD).

2.1 refers to a report around experiences of using Child and Adolescent Mental Health Services (CAMHS) services.

2.2 refers to a report around the care and support for people living with, or affected by dementia.

2. Information and analysis

Both reports can be found in full, including recommendations and service provider/commissioner responses, on the Healthwatch website, or telephone 01773 880786 to request a hard copy.

2.1 CAMHS report (Engagement between May – July 2017)

HWD originally carried out engagement work in 2015 to collect experiences of CAMHS services in Derbyshire. As many of the findings and recommendations in the 2015 report were taken forward by the Futures in Mind Programme it was requested by the Health and Wellbeing Board to repeat a similar engagement to see if changes had had a positive impact.

We spoke to people who had started to use CAMHS within the last 12 months to make sure experiences were current.

Overall, it seemed experiences had improved from 2015 to 2017. Participants gave positive feedback around the flexibility offered to plan sessions at a
convenient time and location. A number of participants explained that the best thing about CAMHS was feeling that help was at hand and that they were now able to talk about how they were feeling.

I would like to highlight the following themes:

- Participants had mixed experiences when talking to their GP about how they were feeling, and explained they would like to see improved training for GPs around mental health, the referral process into CAMHS and the support available.
- Participants told us they would like to see shorter waiting times to begin CAMHS sessions (some participant's had to wait over six months).
- Participants explained they would like to see better information and communication throughout their journey, especially in terms of clear appointment letters explaining who they will see and the role they have so young people know what to expect.
- Many parents and carers felt they had little or no support to help them best look after their child, whilst some appreciated the informal support offered by CAMHS workers.

The HWD engagement officers also reported that many parent/carers did not know about local support groups, even when the information was displayed in CAMHS clinics, they had not taken note of the information.

2.2 Dementia report (Engagement between September 2017 – January 2018)

HWD carried out this piece of engagement work to further explore comments received and enable us to have a greater understanding of the care and support people living with dementia, their carers and family members’ experiences.

The remit was fairly wide and questions were based around the Well Pathway for Dementia.

We spoke to over 100 people.

I would like to highlight the following themes:
Participants highlighted the importance of early education to enable people to become more aware of dementia and understand that it is not a natural part of ageing and that it could happen to anyone.

Most participants felt they received too much information following the diagnosis and felt it would be more beneficial to allow time for the diagnosis to ‘sink in’ and then learn about information and support available. One participant said, “There was loads of leaflets, it seemed bewildering.”

The Derbyshire Dementia Support Service was spoken negatively of in terms of,
1) Poor facilitation of groups with inappropriate activities
2) Lack of trained staff (limited knowledge around dementia)
3) Lack of one-to-one support
- Not all comments were negative, there were a number of people who explained, “They try their best” and “it is getting better.”

Participants spoke positively of the support from Derbyshire County Council homecare staff but highlighted the importance of continuity, routine and training for homecare staff.

Carers felt unsupported in their role in terms of:
1) Not knowing where to go for support (had to become very proactive)
2) Services to support carers were described as ‘disjointed’
3) Caring role having an impact on them being able to look after their own health

Some participants who were self-funding explained their struggles of getting the advice on relevant care and support and felt they were left to “fend for ourselves”.

RECOMMENDATIONS

The Health and Wellbeing Board is asked to note the report themes, and to actively use the report when possible within their own organisations either for wider learning, or in line with an organisational response given to the report where applicable.

Hannah Morton
Engagement Officer
Healthwatch Derbyshire
Child and Adolescent Mental Health Services

Experiences of using CAMHS services in Derbyshire
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1. Thank you

Healthwatch Derbyshire would like to thank all participants and their parents/carers who gave up time to talk to us about their experience of using CAMHS services. We also extend our thanks to the CAMHS services for allowing us to visit their clinics to carry out this engagement activity.

2. Disclaimer

The comments outlined in this report should be taken in the context that they are not representative of all patients, family, friends and carers who have experienced CAMHS services, but nevertheless offer a useful insight. They are the genuine thoughts, feelings and issues that patients, family, friends and carers have conveyed to Healthwatch Derbyshire. The data should be used in conjunction with, and to compliment, other sources of data that are available.

3. About us

Healthwatch Derbyshire is an independent voice for the people of Derbyshire. We are here to listen to the experiences of Derbyshire residents and give them a stronger say in influencing how local health and social care services are provided.

We listen to what people have to say about their experiences of using health and social care services and feed this information through to those responsible for providing the services. We also ensure services are held to account for how they use this feedback to influence the way services are designed and run.

Healthwatch Derbyshire was set up in April 2013 as a result of the Health and Social Care Act 2012, and is part of a network of local Healthwatch organisations covering every local authority across England. The Healthwatch network is supported in its work by Healthwatch England who build a national picture of the issues that matter most to health and social care users and will ensure that this evidence is used to influence those who plan and run services at a national level.

4. Understanding the issue

Healthwatch Derbyshire carried out engagement work in 2015 to collect experiences of using CAMHS services in Derbyshire. Many of the findings and recommendations in the 2015 report have been taken forward by the Futures in Mind Programme, which sets out a vision for children and young people’s mental health in Derbyshire. It was requested by the Derbyshire Health and Wellbeing Board that Healthwatch Derbyshire repeat a similar engagement in 2017, some way in to the delivery of the Futures in Mind Programme, to get up to date feedback from children, young people and their parents/carers about using CAMHS service.
5. What we did in brief

Our engagement team developed a series of questions to ask children, young people and their carers about their experiences of using CAMHS services. Questions covered the whole journey, from first wanting or needing support with mental health, to what it has felt like to use CAMHS services, and if it feels like it has made a difference. We involved people who had started to use CAMHS services within the last year, in order to make sure that experiences were relatively current given the changes triggered by the Futures in Mind Programme. Engagement took place between May-July 2017.

Engagement staff attended a number of CAMHS clinics across Derbyshire to talk to people waiting for their appointments about their experiences.

In the north of Derbyshire, CAMHS services are provided by Chesterfield Royal Hospital NHS Foundation Trust. In this area, we involved 26 participants. Twenty-four people were seen at Chesterfield Royal, one person was seen in the High Peak and one person was seen in Buxton. Fourteen participants were female, 12 were male. The ages of participants were as follows:

- Age 9 - 11 3 no.
- Age 12-14 11 no.
- Age 15-17 12 no.

In the south of Derbyshire, CAMHS services are provided by Derbyshire Healthcare NHS Foundation Trust. In this area, we involved 23 participants. Ten people were seen in south Derbyshire, seven people were seen in Long Eaton and six people were seen in Belper. Seventeen participants were female and six were male. The ages of participants were as follows:

- Age 9 - 11 1 no.
- Age 12-14 2 no.
- Age 15-17 18 no.
- Age not known 2 no.

6. Key findings

- In the first instance, participants talked to either a parent/carer, professional at school or a GP about how they were feeling. Experiences of talking to staff at school were overwhelmingly positive, but were mixed for GPs with a mixture of positive and negative experiences.

- When participants were asked what single thing they would change/improve before seeing CAMHS, several themes emerged:
  - Waiting times to start CAMHS sessions
  - More help, information and support for parents
Improved training for GPs around referrals and support
Clearer appointment letters, explaining who people will see and the role they have so you young people know what to expect
Better information and communication throughout, so that young people and their parent/carers know what will happen, and when it will happen.

- Participants gave generally positive feedback about the flexibility offered to plan sessions in a location that was convenient, and at a time that suited them.
- The overwhelming majority of participants said that the best thing about CAMHS was feeling that help is at hand and that they are talking about how they feel.
- The overwhelming theme from parents/carers was around feeling as if they had little or no support to help them best look after their child.

7. What people told us in North Derbyshire

- Participants were asked about how they were feeling before starting at CAMHS. Responses were a combination of:
  - Depression
  - Anxiety
  - Depression and anxiety
  - Self-harming.

- Participants were asked who they first talked to about how they were feeling, and what happened

Some participants had spoken to a professional at school initially. Participants gave lots of positive comments about their school nurse and/or school counsellor. These relationships felt most useful to participants when support is dependable, and ongoing. Participants had overwhelmingly positive experiences of being referred to CAMHS by their school. Comments included:

- “The school nurse sorted things out quickly and really helped me, I knew she was there if I needed anything.”

- “They seemed to know exactly what to do at school, and they made things happen. It was really good to have somebody that understood.”

Participants gave a more mixed account of the support received and referral process when seeing a GP. Although there are some positive examples, the majority of comments were negative. Example comments include:

- “I talked to my GP but they were not supportive - said it wasn’t their job to do the referral and we would have to speak to a social worker.”
- “The GP didn’t do much.”
- “Wouldn’t refer me, and was very abrupt.”
Participants were asked who had referred them to CAMHS

- GP x 11
- School nurse x 9
- Presented at A&E x 4
- Relate x 1
- Social Worker x 1

Participants were asked how long they waited from being referred to starting sessions

- One month or less x 8
- 1-3 months x 6
- 3-6 months x 10
- 6 months or more x 2

Participants were asked how they felt about how long they had to wait for their first appointment (22 out of 26 participants responded)

- “Quick.” x 6
- “OK.” x 8
- Indicated that a shorter wait, “Would really have helped.” x 3
- “Too long.” x 5

Participants were asked if anything happened to help whilst waiting to be seen by CAMHS

- “Had private counselling.” x 2
- “Support from social worker.” x 2
- “MAT Team/worker helped.” x 2
- “I was seeing the school counsellor.”
- “I could speak to Horizons.”
- “Support from T3.”

Participants were asked if they could change one thing to make CAMHS better for them before starting their appointments, what would it be

- “To be seen more quickly.” x 3
- “More support for parents.” x 2
- “Magazines and a television for when I am anxious waiting”
- “Better training for GPs.”
- “Clearer appointment letters explaining who you will see and the role they have so you know what to expect.”
- **Participants were asked about how convenient the time and location of appointments had been**
  - Participants gave generally positive feedback about the flexibility offered to plan sessions in a location that was convenient, and at a time that suited them.
  - Some adults had struggled to get time off work, and were concerned about children taking time out of school (x 2).
  - Two participants commented that the appointments were a long way from home, with one example being a 60 mile round trip.

- **Participants were asked what is the best bit about coming to CAMHS**
  - The overwhelming majority of participants said that the best thing was feeling that help is at hand and that they are talking about how they feel. People used their own words, such as:
    - “I feel like I can talk to someone.”
    - “I like all of the staff, the receptionists put me at ease when I come in. I can open up.”
    - “Just getting to talk to someone about how I’m feeling, and not bottling it up so I don’t get to a point like I did before.”

- **Participants were asked what is the worst bit about coming to CAMHS**
  - Trying to fit the appointments in with parent’s work x 2.
  - The distance to appointments x 2.
  - Lack of continuity with staff x 2.
  - One parent added that, “At the first appointment, the worker was talking to my child. As a parent it was so hard really hearing how you child is feeling.”
  - “Feeling worried before my appointments.”
  - “Feeling like we aren't getting anywhere.”
  - “I’d like to have only have one person to talk to, I’m happy talking to my CAMHS worker but then my social worker wants to talk to me and the school wants to talk to me and it constantly brings up issues.”

- **Participants were asked if they thought that the CAMHS sessions have helped**
  - The majority of the participants were in the early stages of their appointments, so felt unable to answer this question. However, several people said that they felt as if talking was helping and some spoke about having increased confidence, completing tasks they had previously been unable to manage.

- **Themes raised by parents/carers**
  - **Support**
    - Many parents spoke about feeling as if they had little or no support to help them best look after their child, whilst some appreciated the informal support offered by
CAMHS workers. The Healthwatch engagement staff also reported anecdotally that many parents did not know about local support groups, and even when information was displayed in CAMHS clinics, people had not taken note of this information.

**Negative comments regarding support**

- “I need help in how to support my daughter, there needs to be far more support for parents as I still don’t get any. It is only by chatting to other parents that I find things out.”

- “I feel that the social worker is there to help the child but I don’t get any support as a parent to help me cope with the children and their emotional problems.”

- “I spoke to the school counsellor but this was not at all helpful, as they provided no update to parents.”

- “Parents get no real support.”

- “Any courses are offered during the day. I am a single working parent. Courses need to be held evening time.”

- “I get no feedback as a parent about how my daughter is doing. I need some support; I don’t know how to react with things because I can’t get any confidential information from the CAMHS worker.”

- “No information has ever been given to me about groups.”

- “No signposting information for anything that relates to me as a mum.”

**Positive comments regarding support**

- “I get informal support from my daughter’s worker.”

- “I speak with my child’s workers after the sessions and they give advice and point me in the right direction for support at home.”

**MAT teams**

Several parent/carers spoke about difficulties accessing support from MAT teams, including examples such as:-

- “The MAT support was non-existent.”

- “We had a social worker who was fantastic but she left and we were passed on to MAT worker, who has never been in touch and we have tried to contact her plenty of times.”
8. What people told us in South Derbyshire

- **Participants were asked about how they were feeling before starting at CAMHS. Responses were a combination of:-**
  - Depression
  - Anxiety
  - Self-harming
  - Anger issues
  - OCD
  - Bereaved
  - Behaviour problems
  - Low mood
  - Trouble sleeping

- **Participants were asked who they first talked to about how they were feeling, and what happened**

  The majority of participants had spoken to a parent initially, who then made an appointment to see a GP. A small number of young people spoke to somebody at school first. Participants gave a mixed account of the support received and referral process when seeing a GP. Although there are some positive examples, the majority of the comments were negative. Examples include:

  - “They didn’t really help as they just kept on giving general depression advice such as ‘just go for a walk or do something fun.’ It took about two or three visits before anything actually happened. I think the doctor thought I was just a moody teenager.”
  - “The GP was not supportive, and told my mum that she was not parenting correctly.”

- **Participants were asked who had referred them to CAMHS**
  - GP x 15
  - School nurse x 6
  - Presented at A&E x 2

- **Participants were asked how long they waited from being referred to starting sessions**
  - 1 month or less x 5
  - 1-3 months x 6
  - 3-6 months x 2
  - 6 months or more x 5
  - Unknown x 5
Participants were asked how they felt about how long they had to wait for their first appointment (17 out of 23 participants responded)

- “Quicker than I thought.” x 2
- “OK …” x 5
- “Too long.” x 10

Participants were asked if anything happened to help whilst waiting to be seen

- “I was seeing the school counsellor.” x 3
- “Had private counselling.” x 1
- “MAT Team/worker helped.” x 1
- “GP helped with appointments, and an app.” x 1
- “Support from together for mental wellbeing.” x 1
- “Support from Treetops.” x 1

Participants were asked if they could change one thing to make CAMHS better for them before starting their appointments, what would it be?

- “To be seen more quickly.” x 3
- “Better communication, so you know what will happen next and when it will happen.” x 3
- “More help, information and support for parents.” x 2
- “More information, so you know what to expect.” x 2
- “To be able to ring if it is an emergency.”
- “To be taken more seriously, my referrals kept being refused and I ended up in crisis.”

Participants were asked about how convenient the time and location of appointments had been

Participants gave overwhelmingly positive feedback about the flexibility offered to plan sessions in a location that was convenient, and at a time that suited them. Participants spoke about the service being flexible with early morning, lunchtime and evening appointments to help minimise disruption to school and work.

Participants were asked about what is the best thing about coming to CAMHS

The overwhelming majority of participants said that the best thing was feeling that help is at hand and that they are talking about how they feel. People used their own words, such as:

- “Talking helps.”
- “It is good to be listened to.”
- “I can get advice and help.”
- “Now I can reduce my medication.”
Participants were asked about what is the worst bit about coming to CAMHS

- “Talking about difficult things.” x 3
- “Feeling really nervous about what to expect at the first session.”
- “Cancellations.”
- “Travelling.”
- “Missing school.”
- “Lack of continuity with staff.”
- “Having a long delay to see a psychiatrist.”
- “It feels like a hospital - it could be more colourful and welcoming.”
- “Poor communication between my GP and CAMHS regarding medication.”

Participants were asked if they thought that the CAMHS sessions have helped

Some participants were in the early stages of their appointments, so felt unable to answer this question. However, the majority of people said that they felt as if talking was helping and said that it felt good to be getting the help they felt they needed. One person said that they though they felt better because of their medication, and not because of CAMHS.

Themes raised by parents/carers

Support

Many parents spoke about feeling as if they had little or no support to help them best look after their child, whilst some appreciated the informal support offered by CAMHS workers. The Healthwatch engagement staff also reported anecdotally that many parents did not know about local support groups, and even when information was displayed in CAMHS clinics, people had not taken note of this information. Several parent/carers spoke about the lack of information about community/support groups.

Negative comments regarding support

- “There is a really big gap in support for parents and carers. You do not know what to do for the best. You rely on advice from family but then they are not trained in this sort of thing. Anything would be good, a parent forum, some information, a phone line - just something, as it is hard to cope.”

- “The only support we have had is part of family therapy. We have had no other help and advice on how we as parents and carers cope with things. There was no counselling offered (I have chosen to access this myself privately) and no peer support groups. There was no information booklets or refer to any useful internet help. We did get help from friends who recommended my son’s counsellor.”

- “We had family therapy, but before that you can flounder. For example, there was conflicting guidance about self-harming. One person said to let her self-harm, but to make sure it was done safely and another one said to stop her.
There is a need for parent support group. You need to have the tools as a parent so that you can do the best you can for your child. Parents need guidance.”

- “There is gap in training and support for parents to help parents know how to cope. Parents need mindfulness skills and CBT so they can remain well for their children.”

- “I did parenting a while back, but it was for a wide age range. They showed me videos of younger children sharing things; I needed something for teenagers when they are tearing each other apart. Something more appropriate.”

- “I had a man for family therapy - this was difficult because he was a bloke and so did not work for me.”

- “More help for my mum as she has anxiety issues and me being ill made her worse. She did not get help and did not know what to do.”

- “Got some help through the family therapy but nothing specific for parent/carers and never signposted to anything else or to websites of what to do to help me or my child.”

- “I would like some time with my child’s CAMHS worker to discuss things.”

Positive comments regarding support

- “My daughter’s worker has been a lot more involved and I have been kept in the loop. If I ever have any questions/queries I can always phone and explain and this really helps.”

- “I don’t really get opportunity in the appointments to be involved but I can email or text the worker and they do get back to me. This never used to happen and so this is better than it was before.”

- “I speak with my child’s workers after the sessions which helps me feel as if I have some support when back at home.”

Staff attitude

We received two negative comments about the attitude of staff.

- “At the end of the appointment today we wanted to book a date for another session but the worker said they had left their appointment book upstairs so we couldn’t rebook, which isn’t really good enough.”

- “The receptionists are always bad mouthing other staff members when they are not on shift or in the reception area. This is not very professional.”
9. What should happen now?

- Work to improve the response from GPs to young people and their parents/carers talking to them about mental health, including referrals and support

- Work to minimise waiting times for CAMHS services

- Work to improve the help, information and support for parents throughout the whole process

- Improved information and communication throughout, so that young people and their parent/carers know what will happen, and when it will happen

10. Response from service provider(s)/commissioners

**Commissioners response to Healthwatch Report on CAMHS Services (February 2018)**

Commissioners have found the information in the Healthwatch report very helpful. Additional engagement work is taking place to further consider the points that have been raised, and findings will influence the re-focusing or re-commissioning of service delivery. The report will be considered at the CAMHS contract meetings.

Public Health undertook a robust Health Needs Assessment of children and young people’s mental health and wellbeing across the Derby and Derbyshire area, sponsored by the Future in Mind Delivery Group and published in September 2017. As well as gathering quantitative data from a considerable number of nationally and locally accessible products and services, workshops were conducted as part of the process to capture the qualitative views of children and young people, professionals and other stakeholders, including parents. The resulting product now offers a series of priorities for action in areas including mental illness, vulnerable groups, risk and protective factors, as well as a series of 5 strategic priorities that underpin the refreshed LTP.

Specifically:
- **Strategic Priority 1:** To further develop our engagement and support to parents and carers.
- **Strategic Priority 2:** To develop further a whole-school approach to prevention and early help.
- **Strategic Priority 3:** To develop a new care model responding to children and young people exhibiting complex needs.
- **Strategic Priority 4:** To increase the workforce offer including blended learning approaches across professional groups.
- **Strategic Priority 5:** To develop a place-based approach to interventions and care supporting primary care, developing the voluntary and community sector, linking to schools, and offering digital interventions.

These priority areas will address the, ‘What should happen next statements?’ on p.12 of their report (response from primary care including awareness; minimise waits; supporting parents; improved information and communication throughout. The work on the priority areas will help to mitigate these same issues arising in the future.
In September 2017, the anti-stigma campaign ‘Be a Mate’ was launched. It was developed by an organisation who worked directly with children and young people to ensure that their ideas were incorporated. Posters have been distributed throughout the county and city and a Facebook page has been set up.

One of the key findings from this report has been the lack of adequate information and guidance for parents. Commissioners are now working with the CCG Engagement and Communications Teams to develop a communications strategy to address this.

When participants were asked what single thing they would change/improve before seeing CAMHS, several themes emerged:

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<th>Healthwatch findings</th>
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<td>Waiting times to start CAMHS sessions</td>
<td>Commissioners are working with both CAMHS providers to reduce waiting times. North CAMHS have done a detailed analysis of why they are struggling to achieve reasonable waiting times. The CCG is considering national research to assess whether there is sufficient funding per head of population to achieve low waits. Commissioners are considering whether or not it would be appropriate to include a maximum wait time of 12 weeks to the first appointment, for all contracts relating to emotional health and wellbeing, and asking for an exception report for each case where this is not realised. Given the increase in demand, so much of the discussion needs to focus on how to manage demand within the community, and ensure that those children that do need a service are signposted to appropriate provision as soon as possible, and do not have to wait too long to be seen. SPOA is used in the south of the county to triage referrals to CAMHS. In the north and Erewash, a community based triage service has now been commissioned to ensure that referrals to CAMHS which might be best responded to by another organisation eg. Relate, School nursing service, MAT Team, Action for Children etc. are responded to by the appropriate organisation in a timely manner.</td>
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<td>More help, information and support for parents</td>
<td>Parents support has been commissioned for 2018/19, and will be reviewed during the</td>
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From 2019 there will be an STP footprint-wide offer for parents. This will include awareness raising, how to identify signs of early problems, how to provide early support to reduce the likelihood of problem escalation, how to recognise that your child needs additional support and how to get that additional support.

During 2018/19 there will also be a review and consideration of the support which may be commissioned for specific cohorts of parents/carers, such as foster carers.

Commissioners will include information and advice to parents about how to support their child’s mental and emotional wellbeing as a requirement in any future commissioning of CAMHS.

Commissioners have started to develop a communications strategy and this will be included in that.

| Improved training for GPs around referrals and support | Commissioners will work with primary care colleagues to improve the training and information for GPs.  
As ‘Place’ develops, the commissioners will ensure that GPs are aware of the provision in their area and will develop simple ways for GPs to refer in to the appropriate provision.  
During 2018/19 there will be a review of the points of access that are used for referrals, with the aim of having a much clearer system for GP referrals to ensure that the child receives the right service as soon as possible.  |
|---------------------------------------------------------------------------------------------------------------|
| Clearer appointment letters, explaining who people will see and the role they have so young people know what to expect | Commissioners have started to develop a communications strategy and this will be included in that.  
Commissioners will include this as a requirement in any future CAMHS contract.  
CAMHS will be required to consult with their service users and with parents/carers on a regular basis to inform continuous service improvement.  |
| Better information and communication | Commissioners will consider how young |
throughout, so that young people and their parent/carers know what will happen, and when it will happen.

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<th>Participants gave generally positive feedback about the flexibility offered to plan sessions in a location that was convenient, and at a time that suited them.</th>
<th>Flexibility in appointments, as close to home as possible will be included in any future commissioning of CAMHS.</th>
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<th>The overwhelming theme from parents/carers was around feeling as if they had little or no support to help them best look after their child.</th>
<th>CAMHS services in both north and south of the county now have their own websites that include sections for parents. Commissioners will require CAMHS to further develop guidance for parents on those websites.</th>
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<th>Some adults had struggled to get time off work, and were concerned about children taking time out of school.</th>
<th>Commissioners will consider whether there should be a change in the times that CAMHS is commissioned to work, and whether they should be commissioned to provide any evening appointments.</th>
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<th>I’d like to have only have one person to talk to, I’m happy talking to my CAMHS worker but then my social worker wants to talk to me and the school wants to talk to me and it constantly brings up issues.</th>
<th>CAMHS will be expected to ask if the young person if s/he is happy for their information to be shared as this would reduce the number of times a story has to be repeated. If so CAMHS will obtain written consent and will share information with others as requested. CAMHS in the north are now using System One which should assist with information sharing if this is requested. This will bring a more consistent approach across Derbyshire.</th>
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<th>The Healthwatch engagement staff also</th>
<th>Commissioners will do further engagement.</th>
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</thead>
</table>
reported anecdotally that many parents did not know about local support groups, and even when information was displayed in CAMHS clinics, people had not taken note of this information.

 Commissioners have started to develop a communications strategy and this will be included in that.

<table>
<thead>
<tr>
<th>Any courses are offered during the day. I am a single working parent. Courses need to be held evening time. There is a need for parent support group. You need to have the tools as a parent so that you can do the best you can for your child. Parents need guidance.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Commissioners will consider commissioning information sessions/workshops for parents in the evenings. The commissioners will ask the CVSs to support any self-help groups for parents that arise from the workshops/training.</td>
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<table>
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<tr>
<th>To be able to ring if it is an emergency</th>
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<tbody>
<tr>
<td>Commissioners will work with CAMHS to ensure that there is clear and consistent information for children and young people and parents/carers on how to get help in an emergency and information is on respective websites.</td>
</tr>
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**Derbyshire Healthcare NHS Foundation Trust Response**

**Parent/carer engagement**

The CAMHS service is committed to developing participation at all levels, we have recruited a young person to the role of Expert by Experience, and have an identified participation lead to support them. We are currently reviewing our original plan to recruit a Parent Expert by Experience to ensure we can capture a wide range of views.

The service is continuing to develop the information that is provided to young people their families and carers, work also continues in the development of the CAMHS website. This includes information about services, how to access services, frequently asked questions, and useful links to external resources. The service information leaflets will be reviewed annually with oversight by the expert by experience workers to ensure that leaflets are clear, informative and easy to read. Work to undertake this review of leaflets is being planned and will commence in April 2018.

A Parent Support Group is being established and will recommence from Temple House, Derby and Rivermead, Belper by April 2018. We will provide details of this in the waiting rooms and also on our website.

A waiting list initiative has been established and embedded into current practice to ensure that young people and their families who are waiting for therapy are reviewed by the duty worker and are aware of how to access support should their circumstances change. Urgent
face to face review appointments are available if required. Initial feedback of making contact in this way is positive.

Finally, the waiting rooms within CAMHS services have been reviewed by young people and parents and a key initiative is to improve the environment and the information available. We value feedback on the service we provide to support quality improvement. Our waiting rooms provide opportunities to provide feedback, and include visual initiatives such as “You Said, We Did”. We are seeking the view of young people and carers to help select new furnishings in Temple House at present.

Since reading this report, we have already made the following changes:

We have introduced quality improvement project which is an additional waiting list initiative, including the additional investment of time for staff to increase the level of frequency in contacting young people, parent/carers waiting for a CAMHS assessment to check if an urgent appointment is required, and to advise about what other support services are within their area to help, support and provide advice to parents/carers. This approach commenced in January 2018.

The waiting list initiative is supported by an additional urgent review appointment slot which is available to support families/young people waiting to be seen if required. We have developed an Early Access pathway which means that there is one streamlined waiting list for CAMHS, which improves equality of access across the South Derbyshire rather than by locality (Amber Valley, Erewash, South Derbyshire, Derby City). This means that young people referred can access all professionals in their area rather than having local blockages in waiting time. As a result of this change we have an average wait time now of 9.15 weeks, and we continue to monitor our waiting times. We will continue our work to review how we work and what we do, in order that we can continually improve our responsiveness and reduce the time to access our services.

We are embedding a parent participation group across the whole service which is working on themes which include: inclusion, communication, and “You Said, We Did.” We hope this goes some way to improving how we involve parents.

We have collaboratively developed (with parents/young people) a CAMHS website to improve access to information for young people, parents/carers and professionals. This is important to share information on what to expect and learn from feedback. Whenever we gain feedback from families on things we can improve, we will consider our information on our website so we can improve what information we share and how we do for everyone.

Parents and young people have reviewed all waiting areas and started to improve information available and the environment. Our Young People’s Participation Group are working with us to improve the furnishing, décor and activity in our waiting areas which we will undertake during 2018/2019.

b) We will also be making the following changes:
<table>
<thead>
<tr>
<th>Key Themes</th>
<th>Milestones/Objectives</th>
<th>Actions required</th>
<th>Lead</th>
<th>By when</th>
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</thead>
<tbody>
<tr>
<td>Waiting times to start CAMHS sessions</td>
<td>Recruitment to vacant posts</td>
<td>Advertise positions and recruitment is in process</td>
<td>Operations Managers and Clinical Leads</td>
<td>Commenced December 2017. February 2018</td>
</tr>
<tr>
<td></td>
<td>Improve access to evidence based interventions</td>
<td>Development of a NVR parent intervention programme</td>
<td></td>
<td>Monthly monitoring by Area Service Manager of waiting times. Shared with Commissioners. Mechanism in place.</td>
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<tr>
<td></td>
<td>Improve access to CAMHS</td>
<td>Reduce waiting times from referral to first/second appointment</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>Continued support to the Future in Mind programme improving early access to evidence based interventions throughout children and young people’s services.</td>
<td></td>
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<tr>
<td>Support to families who are waiting for CAMHS.</td>
<td>Embed waiting list initiative as core practice in CAMHS</td>
<td>Review and monitor the waiting list initiative</td>
<td>Operations Managers and Clinical Leads</td>
<td>Commenced January 2018. Review March 2018.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Provide clear information about who the families are able to contact (CAMHS duty - routine, CAMHS RISE - urgent)</td>
<td></td>
<td>February 2018</td>
</tr>
<tr>
<td>More help, information and support for parents</td>
<td>Parent Participation Groups Parent support groups in secondary care services Web based information Parent Expert by Experience</td>
<td>Continue to work on the CAMHS web site Continue to develop the information that we provide to young people/parent/carers. Information leaflets regards specific</td>
<td>Operations Managers and Clinical Leads and Expert by Experience</td>
<td>Commenced January 2018</td>
</tr>
</tbody>
</table>
| Improved training for GPs around referrals and support | Regular attendance at GP training forums  
Work with Commissioners to support the delivery in primary care | Develop a training / information pack for GP’s. Encourage GP’s to utilise Duty system for any queries. Review roles of PMHW’s/SCA’s to improve GP’s access to specialist advice. | Operations Managers and Clinical Leads and medical team. Oversight by ASM. | Develop programme by March 2018.  
Six months |
| Clearer appointment letters, explaining who people will see and the role they have so young people know what to expect | Information leaflet to be sent with every initial / follow up appointment | Link to the CAMHS website and Local Offer  
Review information leaflets annually or after any significant change | Operations Managers and Expert by Experience Workers | Starting January 2018  
Annually or as required |
| Better information and communication throughout, so that young people and their parent/carers know what will happen, and when it will happen | Collaborative care plans - signed by y/p / parent/ carer  
E information available to young people/parents and carers | Improve CAMHS website  
Support the development of the website to become mobile friendly  
Ensure that the right information is available at the right time for the right people - review admin/letter.  
Audit care plans across CAMHS  
Consider creative use of social media | Operations Managers and Clinical Leads | Starting January 2018  
Care plan audit planned for February 2018.  
Six months |
| Feedback on staff attitude and conduct | Standard of high levels of communication and being able to leave an appointment knowing what comes next.  
Feedback to staff | Set clinical and operational standard of young person leaving an appointment knowing exactly what happens next and when.  
ASM to meet with all | ASM CAMHS | March 2018 |
<p>| around customer care standards. | admin staff to feedback findings. Identify any training needs and source if required. | March 2018 |</p>
<table>
<thead>
<tr>
<th>Item of Concern</th>
<th>Response</th>
<th>Timescale</th>
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<tbody>
<tr>
<td>Waiting times for appointments and continuity of staff</td>
<td>CAMHS is working on improving the written information that is available for families whilst waiting (see below). CAMHS is currently collaborating with Commissioners to work on reducing waiting times and improving continuity for CAMHS services.</td>
<td>April 2018 July 2018</td>
</tr>
<tr>
<td>Information and support for parents</td>
<td>All clinicians have been reminded of the importance of keeping parents informed of CAMHS work, as far as confidentiality and consent allows. This has been discussed at Clinical Leadership Team meetings and cascaded via team meetings. Feedback will continue to be monitored in relation to this point. CAMHS is currently developing a dedicated website, which will act as a means of sharing general information with all service users, families and carers. The comments provided by Healthwatch within this report will be taken into account when designing the website.</td>
<td>Complete April 2018</td>
</tr>
<tr>
<td>Clearer information regarding what to expect</td>
<td>CAMHS is in the process of reviewing information sent out to families with a first appointment. Currently, a national CAMHS leaflet is provided; the team wish to produce a bespoke information leaflet that is more relevant locally and includes local contact details. It is expected that this will support service users and alleviate any associated anxieties; service user feedback will be regularly reviewed to monitor any improvements. CAMHS also anticipate that the new website will provide a useful means of sharing information about the service and what can be expected.</td>
<td>April 2018</td>
</tr>
<tr>
<td>Training for GPs</td>
<td>In mid-2017, a new team of Specialist Community Advisors was implemented in CAMHS; there are 5 advisors across North Derbyshire who are currently developing links with colleagues in the community, including GPs. The role involves consultation, advice and training; the intention is that this work will support people in making contact with CAMHS.</td>
<td>Complete</td>
</tr>
<tr>
<td>Availability of magazines and a television whilst waiting</td>
<td>In 2017, CAMHS engaged with a team of young people, via a college scheme, who helped to revamp the CAMHS waiting area at The Den. The young people designed some decorative features for public areas and the waiting area is now equipped with a television and relevant reading materials.</td>
<td>Complete</td>
</tr>
<tr>
<td>Distance to appointments and flexibility of times</td>
<td>CAMHS do try to offer flexibility as much as possible, with bases at Buxton, Chesterfield and Bakewell; in addition CAMHS staff are able to access rooms in some schools. Some teams do offer out of hours appointments and CAMHS are currently exploring options with Commissioners for expanding this further.</td>
<td>July 2018</td>
</tr>
</tbody>
</table>

11. Your feedback
Experiences of using CAMHS services in Derbyshire

Healthwatch Derbyshire is keen to find out how useful this report has been to you, and/or your organisation, in further developing your service. Please provide feedback as below, or via email.

1) I/we found this report to be: Useful / Not Useful

2) Why do you think this?

......................................................................................................................................................................
......................................................................................................................................................................
......................................................................................................................................................................

3) Since reading this report:

a) We have already made the following changes: ..............................................................
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b) We will be making the following changes: ..............................................................
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Your name: ...........................................................................................................................

Organisation: ...........................................................................................................................

Email: .................................................................................................................................

Tel No: .................................................................................................................................

Please email to: karen@healthwatchderbyshire.co.uk or post to FREEPOST RTEE-RGYU-EUCK, Healthwatch Derbyshire, Suite 14 Riverside Business Centre, Foundry Lane, Milford, Belper, Derbyshire DE56 0RN.

V4 CAMHS report 2018 24042018 HH-S
Dementia Report

Experiences of Health and Social Care services in Derbyshire from the perspectives of people living with dementia, their carers and family members

Date: March 2018
Author: Hannah Morton
Job Title: Engagement Officer
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</table>
1. Thank you

Healthwatch Derbyshire would like to thank all participants who gave their time to talk to us about their experiences of health and social care service in Derbyshire in relation to dementia care. We also extend our thanks to the many groups and services who supported and cooperated with this engagement activity.

2. Disclaimer

The comments outlined in this report should be taken in the context that they are not representative of all patients, family, friends and carers who have experienced health and social care service in relation to dementia care but nevertheless offer a useful insight. It is important to note that the engagement was carried out within a specific time frame and therefore only provides a snapshot of patient experience collected at that point in time. They are the genuine thoughts, feelings and issues that patients, families, friends and carers have conveyed to Healthwatch Derbyshire.

The data should be used in conjunction with, and to complement, other sources of data that are available.

3. Background

Healthwatch Derbyshire is an independent voice for the people of Derbyshire. We are here to listen to the experiences of Derbyshire residents and give them a stronger say in influencing how local health and social care services are provided.

We listen to what people have to say about their experiences of using health and social care services and feed this information through to those responsible for providing the services. We also ensure services are held to account for how they use this feedback to influence the way services are designed and run.

Healthwatch Derbyshire was set up in April 2013 as a result of the Health and Social Care Act 2012, and is part of a network of local Healthwatch organisations covering every local authority across England. The Healthwatch network is supported in its work by Healthwatch England who build a national picture of the issues that matter most to health and social care users and will ensure that this evidence is used to influence those who plan and run services at a national level.

3.1 What is dementia?

According to the Alzheimer’s Society, “the word ‘dementia’ describes a set of symptoms that may include memory loss and difficulties with thinking, problem-solving or language, and can often cause changes in mood, perception or behaviour. Dementia is caused when the brain is damaged by diseases, such as Alzheimer’s disease or a series of strokes. The specific symptoms that someone with dementia may experience will depend on the parts of the brain that are damaged and the disease that is causing the dementia”.

V1 Dementia Report 22/02/2018 HM
Types of dementia may include:

- **Alzheimer’s disease:** This is the most common form of dementia; this is when abnormal proteins cause brain cells to die and early symptoms are often associated with memory loss.

- **Vascular dementia:** The second most common type of dementia; this can occur when the blood supply to the brain is interrupted, for example, after a stroke, and causes problems with thinking and planning.

- **Dementia with Lewy Bodies:** This type of dementia gets its name from the tiny spherical structures that develop inside nerve cells. Symptoms often include fluctuating alertness, hallucinations and problems with movement.

- **Frontotemporal dementia:** Damage occurs first in the front and sides of the brain; personality, behaviour or language are initially more affected than memory.

- **Young-onset dementia:** People who are diagnosed with dementia before they are 65 are said to have ‘young-onset dementia.’ They may have different challenges to older people, partly due to their age.

For more information, visit the Alzheimer’s Society: [https://www.alzheimers.org.uk/info/20007/types_of_dementia/1/what_is_dementia](https://www.alzheimers.org.uk/info/20007/types_of_dementia/1/what_is_dementia)

### 4. Rationale for the report

To ensure a diverse range of individuals are able to share their views on local health and social care services, Healthwatch Derbyshire undertake targeted pieces of work, paying specific attention to those who may otherwise struggle to be heard. The Intelligence, Insight and Action Sub Group of Healthwatch Derbyshire, who regularly appraise all the comments and experiences received by the organisation, recommended this engagement priority to further explore comments received by Healthwatch Derbyshire and enable Healthwatch to have a greater understanding of the care and support people living with dementia (PLWD) and their carers/family members experience.

### 5. What we did in brief

To collect consistent information across the county, a series of questions were developed to provide a framework for discussions. These were based around the ‘Well Pathway for Dementia’ (see appendix 1, p34) which was developed by NHS England who are working towards ensuring people diagnosed with dementia by 2020 have a better experience of health and social care support, from diagnosis through to the end of life. The ‘Well Pathway for Dementia’ provides a framework to align and co-ordinate the contribution of health and social care partners to meet key commitments which include: improving diagnosis, support and care after diagnosis, enabling people to live well in their own homes for longer, end of life care and education, training and workforce. More information can be found: [https://www.england.nhs.uk/mental-health/dementia/](https://www.england.nhs.uk/mental-health/dementia/)
5.1 Methods of engagement:

All engagement officers (EOs) undertook a Dementia Friend’s training session prior to engagement starting to ensure a good understanding of dementia.

We collected people’s experiences in a number of different ways over a four-month period (September 1st 2017 - January 19th 2018):

- **One-to-one interviews:** We initially visited different groups and services to establish contact with PLWD and their carers and/or family members. A date was then set with these individuals for a one-to-one interview to take place. A total of 50 interviews took place
- **Group discussions:** The EOs visited and facilitated discussions at the Derby and Chesterfield Dementia Engagement and Empowerment Project (DEEP) groups, the Working Age Group and a Black Minority Ethnic (BME) Forum. Approximately 47 participants took part in these discussions
- **Dementia-Friendly survey:** We produced a short survey containing four questions to capture experiences directly from PLWD. This was completed by 13 PLWD
- **Professionals’ comments:** We welcomed feedback from professionals.

The chart below shows the composition of the participants:

![Chart showing the composition of participants](chart.png)

Some of the feedback was provided by PLWD independently, whereas some feedback was provided jointly by the PLWD and their carers and some feedback was from family members who did not class themselves as carers.

A number of considerations should be made when reading the findings, which include:

- Some participants struggled to recall all elements of their experience
- To ensure confidentiality we have removed information that we feel could potentially lead to the identification of any of the participants
- Some of the experiences date back over 12 months ago, however all participants were currently accessing services at the point of engagement.

6. Key findings

Preventing well:

- Participants highlighted the importance of early education to raise awareness and understanding of dementia
- There was an expressed need for more information around all types of dementia, not just Alzheimer’s disease which seems to be the main focus
- A high number of participants discussed how they managed as a family for a long period of time without help, as they were unsure of the signs and symptoms
- Participants felt it was important that all the signs and symptoms were communicated, rather than focusing on just memory loss, as this would make it easier for people to recognise the signs and seek professional advice at an earlier stage
- Participants under the age of 65 felt that it was difficult to access information about young-onset dementia. It was also felt that images used for literature and campaigns portrayed a negative image, and sent out false messages that dementia only affects older people.

Diagnosing well:

- A high proportion of participants felt their initial concerns were listened to by a professional and a referral was made to a Memory Assessment Clinic (MAS Clinic) for further investigation. However, some carers and family members felt professionals did not always listen to the concerns they expressed around their loved ones, which caused a delay in being referred and ultimately receiving the diagnosis
- Some participants under the age of 65 felt their symptoms were sometimes “downplayed”
- Most participants felt their diagnosis was delivered in a compassionate manner, but some described it as “very matter of fact”
- Most participants received their diagnosis within a few months, and those who waited longer were being monitored during this time
- The BME community felt that they had experienced longer waits for a formal diagnosis due to language barriers and cultural differences
- A high proportion of participants felt they received too much information at the point of diagnosis and felt it would be more beneficial to have information around their diagnosis once it had “sunk in”
- A number of carers and family members felt there was no opportunity to ask questions privately about their loved one’s diagnosis
- Not all participants were referred/signposted to sources of support following diagnosis. However, those who had been referred/signposted, had mixed experiences of the support received
Most participants explained they did not have a care plan in place, or were unsure whether one had been developed. Some participants did have a care plan in place with Derbyshire County Council (DCC) or were linked with the Derbyshire Dementia Support Service (DDSS).

The experiences of reviews within the first year varied; some participants had a yearly review, whereas some participants had no update of their diagnosis.

Supporting well:

- There were mixed experiences of accessing support from health and social care services, with many highlighting difficulties.
- Participants who were linked in with their GP or a health professional spoke positively of the support received.
- Experiences of dementia key workers and ‘dementia-specific’ wards within hospital settings were positive.
- There appeared to be language barriers especially within hospital settings for participants from the BME community.
- There was a perception that some professionals working with PLWD in a wide range of services had not been adequately trained to meet their care and support needs.
- In terms of care at home, experiences were mixed depending on the service provider. Participants who used care services from DCC spoke positively about the care received and the level of training from the staff.
- Participants highlighted the importance of continuity, routine and training for homecare staff.
- Information and support from DCC and occupational therapists (OTs) was valued by participants with regard to falls prevention and home adaptions.
- Participants appreciated the Safe and Well checks offered by Derbyshire Fire and Rescue Service (Derbyshire FRS), which provided reassurance.
- The support/activities provided by independent groups and day centres were spoken highly of by all.
- The DEEP groups were praised by participants.
- DDSS was spoken about negatively in terms of participants feeling that there was poor facilitation, inappropriate activities and a lack of one-to-one support. Participants explained they would prefer the groups to be a place of meaningful support with activities that are stimulating and educational. Some participants did share positive experiences, and others felt that the service was “getting better”.
- Participants explained they would prefer to have a named person to go to for advice and support as there is often ‘too many’ organisations that offer support which makes the process of ‘knowing who to go to’ fairly confusing.
- Participants under the age of 65 felt there was a lack of support and activities available for young-onset dementia.

Living well:

- Most participants identified themselves as carers, however there were mixed experiences of having a carer’s assessment.
Carers spoke of initially being unsure of where to go for support and therefore had to become very proactive in finding out what was available to them.

The co-ordination of services to support carers was described as ‘disjointed’.

Participants who were self-funded explained they were often just given a booklet or leaflet on the support available with no further help and often felt they were left to “fend for themselves”.

Participants felt unsupported in their role as a carer and some participants explained their caring role had an impact on them being able to look after their own health.

Responses were mixed regarding the carer’s emergency card, however those who were not aware of the card explained they would have found it useful.

Most participants who had registered as a carer with their GP surgery spoke positively of the support received.

There seemed to be a lengthy process in accessing respite care, however once the support was in place it was spoken positively of and valued by carers.

Experiences related to Continuing Healthcare (CHC) were poor, with participants explaining the process is difficult to understand.

Participants mainly spoke positively around Dementia Friendly Communities (DFC), with most participants explaining ‘it is getting better’.

Experiences around Dementia Friendly Swimming sessions and Dementia Friendly Walking were all positive.

7. What people told us

Experiences of people living with dementia over the age of 65

7.1: Preventing well

Early education:

Participants felt that early education is important to enable people to become more aware of dementia and understand that it is not a natural part of ageing and that it could happen to anyone.

Participants felt that there was not enough information around risk reduction and the early signs of dementia. Some said that “it is getting better” however, others explained that “if it isn’t something that is affecting you, you don’t really take note or talk about it”.

Sample of comments:

- “I don’t think people link health and dementia together; I feel people believe there is nothing you can do”
- “I think people believe it is part of getting old”
- “I think people are frightened and therefore do not particularly want to look at what they could do to prevent dementia”.

V1 Dementia Report 22/02/2018 HM
Information around all types of dementia and the different symptoms:

Participants felt there should be more awareness around all types of dementia, including the different signs and symptoms.

It was felt that there is a lot of awareness raising around Alzheimer’s disease, and although it is understood that this may be the most common form of dementia, participants felt there was not as much information available for the other forms, such as vascular dementia and frontotemporal dementia.

Participants felt there should be more information around the different signs and symptoms as many people believe it is solely around memory loss which many participants explained that this was not their first sign.

The majority of the participants firstly noticed signs which included difficulty completing DIY tasks, withdrawal from social activities, changes in behaviour and becoming lost in familiar surroundings.

Sample of comments:

- “There is not enough information on all types of dementia; it all seems to be around Alzheimer’s. For my mum it was not around memory loss, she had frontotemporal dementia where the signs were around speech and randomised comments but we were not aware of this at the time”
- “It is not always about memory loss, it can be mood changes and putting things in order as it was for my husband”
- “If people knew more about the signs they would be able to seek help and get diagnosed more quickly, but I think it is also about having the confidence of knowing what to do”
- “He had always been good with DIY but he began to struggle with this and had difficulties with problem-solving and was becoming very frustrated at things”
- “I suppose it was the memory loss but this was not the main issue; it was his behaviour and getting lost”
- “I think people should be thinking about symptoms other than memory loss”
- “Maybe if there was better awareness you would go to your GP sooner and it would not be so dragged out and feel so long”
- “I hesitated about seeking medical help due to conflicting symptoms of depression”.

7.2: Diagnosing well

➢ Before the diagnosis

Unsure of the early symptoms and when to seek professional advice:

As reflected in the preventing well section of the report, many participants felt there should be more information around other symptoms of dementia, rather than a focus on
memory loss.

Once the signs had been recognised by the individual, family or carers, quite a few participants explained they had “discussed it as a family” with many continuing to “manage as a family” for a lengthy period of time before seeking professional advice. As some participants explained, “it was a sore subject to bring up” and “it was just little things to start with”.

Most participants visited their GP quite quickly or at least within the first year of noticing the signs. Those who waited much longer explained this was due to “being unsure of the signs” and in a few cases they had put it down to age, stress or depression.

Initial discussion with a professional (mainly a GP):

A high proportion of participants did feel their initial concerns were listened to. Most GPs carried out a Mini Mental State Examination (MMSE) and then made a referral to the Memory Assessment Service (MAS) clinic, whereas some participants were told to see how things go and monitor any changes.

Sample of comments:

- “Yes, they arranged for the usual tests; they did a paper-based cognitive test at the surgery and checked for urinary tract infections and vitamin deficiencies as this can eliminate other things”
- “Very much so, all the staff I encountered pre-diagnosis were excellent and very thorough”
- “I did not feel my concerns were taken seriously initially by the GP as it was put down to old age and it wasn’t until we returned to the GP with the same concerns that a referral was made”
- “We had to wait 18 months before anything was acted on. You shouldn’t have to keep going back and have to ask for a referral. There should be continuity with GPs too; we saw a different doctor each time”
- “We went to the GP who did a test with items on a tray. We were told nothing was wrong as he was in the normal range”.

The views of carers and family members:

Some carers and family members felt professionals did not always listen to the concerns they expressed around their loved ones, which caused a delay in being referred and ultimately receiving the diagnosis.

Sample of comments:

- “Professionals should listen to the family as we are with them and can see the changes properly. When people go to their doctors they could be at their ‘best’ and the doctors may not always see the person when they are manifesting dementia symptoms”
- “We went to the GP who did a memory test and he sailed through it. It was then put down to old age, but when you have lived with someone for so long you know when something isn’t right. We left it for a few months, we ignored it and hoped it would go away”
- “When pushing for doctors to listen, you have to trust your instincts as you know when there is something wrong with your husband or wife. You should not have to keep putting your foot down to get a response and a diagnosis”.

➢ Receiving the diagnosis

Referral to the MAS clinics:

Most participants explained they underwent various tests and scans with a specialist, including computed tomography scans (CT scans) and magnetic resonance imaging scans (MRI scans) to determine their diagnosis.

Most participants felt their diagnosis was delivered in a compassionate manner, whereas some participants expressed a negative opinion of how the diagnosis was delivered.

Sample of comments:

- “The consultant seemed very positive and caring”
- “I felt the delivery was tailored to suit us”
- “It was very cut and dry; this is dementia and you have to deal with it”
- “It was just a matter of fact for the doctor but for us it was devastation. They need to think of this. When my mother had cancer all the professionals were extremely compassionate and understanding. Why is this not always the case with dementia?”

Length of time from referral to receiving a diagnosis:

Most participants felt that this happened quite quickly with the majority of participants receiving their diagnosis within a few months. Some participants had to wait much longer. But most of these participants explained that they were under a specialist and were undergoing various tests.

Sample of comments:

- Most participants explained that “it happened quite quickly”
- “My diagnosis was completed in 10 months; there was a lot of confusion over the scans and the wait just felt too long”.

Information following diagnosis:

Most participants felt they received too much information following the diagnosis and it was felt that it would be more beneficial to allow the diagnosis to “sink in” first, and then have the opportunity to receive information around the diagnosis and the support available.
Many participants did not have concerns immediately after diagnosis and felt they needed time to “come to terms with the diagnosis”. Some carers/family members who did have immediate concerns felt there was no opportunity to ask questions privately away from their loved ones.

Some participants had received no information regarding their diagnosis and a few were unaware of the type of dementia and had undertaken their own research. It was felt by some participants that there was no clear answer to the questions they had around progression of the disease.

Sample of comments:

- “It was like a lead balloon; it was such a shock and at that point we found it very difficult to digest any information that would be relevant to us later”
- “You are given too much information covering everything. It would be useful to receive only the information that you need, to make it easier to take in and understand”
- “There was loads of leaflets, it seemed bewildering at first but I know the information is somewhere; it is just knowing what leaflet to look at first to find what I need”
- “It would be best to have a follow-up a few weeks after the diagnosis to be able to ask questions and understand everything better”
- “I was a bit shell-shocked, so I didn’t ask any questions at the time but then I never got the chance to ask my questions because we didn’t see the consultant again”
- “I wanted to ask questions but I felt this was inappropriate in front of my father. I felt there should have been another opportunity given for family members to ask questions”
- “It was very difficult to ask questions in front of mum when she was still in denial”
- “There was no clear answer to the questions we asked such as the progression, understandably they cannot give the exact date as everyone is different but an estimate would have been good, as this has affected our life insurance”.

The feelings of people following diagnosis are documented below:
Signposting to support following diagnosis:

Not all participants were referred to further support following their diagnosis. Some participants who were referred found it useful.

Some participants were referred to the Living Well Programme, although this was not consistent for all participants. Those that attended the programme spoke highly of it and explained how informative they found the sessions.

Some participants (mainly in Erewash and South Derbyshire) explained the Q&A sessions organised through Derbyshire Healthcare NHS Foundation Trust (DHcFT) were very helpful.

Sample of comments:

- “No one was helping us come to terms with the diagnosis; it has felt like a bereavement”
- “We thought getting the diagnosis would help us; the MAS clinic became this magical place where we would get the information and help we needed, but this was not the case”
- “There is a lack of professionals involved post-diagnosis. I have contact details for a community psychiatric nurse (CPN) and dementia nurse but you have to chase it up yourself rather than them contacting you which would have been better”
- “Straight after the diagnosis the consultant took us to meet someone from Making Space and we were then told all the facts”
- “It was nice to meet people who are going through the same as you; we also learned a lot and received a lot of information” (Living Well Programme)
- “It brought both carers and PLWD together and we were in separate rooms which allowed us to share experiences without the individuals getting upset when discussing certain subjects” (Living Well Programme)
- “They give you a chance to ask questions” (Q&A sessions DHcFT)
- “They are good as you can share tips with other carers” (Q&A sessions DHcFT).

Care Plans:

The majority of participants responded “no” or “unsure” as to whether they had a care plan in place following their diagnosis. A small number of participants said they had a care plan with DCC and others said they had been linked to the DDSS.

Sample of comments:

- “I believe a care plan was developed early on, but nothing has ever come of it”
- “We had a home visit from Making Space and the dementia advisor explained things around a care plan but I am unsure whether it has been developed or not”.

Review within first year of diagnosis:
Some participants were unable to provide an answer due to being recently diagnosed. Some participants talked about a yearly review, whilst others said that they had no review or update on their diagnosis at all.

A number of participants explained they were under the Community Mental Health Team (CMHT), doctor or consultant and felt their diagnosis was being monitored.

Sample of comments:

- “We have a review booked with the consultant, but this is 18 months after the diagnosis and there has been nothing in between”
- “We go every year for a review, but I think they are a waste of time as my wife can no longer read and she does not understand what is happening. It is not a pleasant experience for us”.

7.3: Supporting well

Support from Health and Social Care services:

There were mixed experiences of support following diagnosis. Some participants did not feel they needed support as they were “coping fine at the moment”. A number of participants explained they now have the support in place but it has taken a long time.

Some GPs were seen to be very supportive. A few participants were still under their consultants although as mentioned previously not all participants had regular appointments/reviews.

When asked if participants had a ‘named support worker’ the majority did not have one. Others talked about their dementia advisor (DA) from Making Space, their CPN, social worker or care co-ordinator.

Sample of comments:

- “We do now have the support in place but it has taken so long as I have had to do all the organising”
- “It has been a long, painful, emotional and stressful journey along the dementia pathway so far but we have been shown so much love and affection, it has been overwhelming”
- “It has felt like hard work; as a family we have had to be proactive in getting the information and the support we needed”
- “He (the GP) has been excellent and attentive and has been honest with us about my mum’s condition worsening”
- “We were put in touch with our care co-ordinator and she was marvellous”
- “We thought we had a named social worker, until we contacted them and we were told we had been discharged from the service as we had not been in contact for a while”
- “My friend has recently been diagnosed with breast cancer. All the support was put in place for her straight away, however when you are diagnosed with dementia, if you are seen to be managing and coping OK you don’t get any help at all”
- “People living with dementia are not always taken seriously. The system is not geared up for conditions such as dementia because you cannot see it. If my mum had cancer, the support would have been put in place for her”
- “There needs to be better communication between services. I have only recently been notified of the financial support and what is available to us and I feel I should have been notified sooner so I could have planned things better”.

Experience of hospital:

There were mixed experiences on hospital wards. One participant explained that dementia key workers are not available on all wards. Participants said that on the whole, once they had explained that their loved one had dementia the quality of care seemed to improve.

Many participants explained how crucially important it is for professionals to explain what they are doing.

Not all staff knew about ‘This is Me’: 
https://www.alzheimers.org.uk/info/20033/publications_and_factsheets/680/this_is_me

Sample of comments:

- “The staff are excellent on dementia-specific wards, but on others there is a lack of awareness and training so staff may think someone is being odd or aggressive when in fact, it is the dementia causing them to act this way”
- “My husband went into hospital with an injured leg. The staff were not aware that he had dementia or how to deal with him and I feel he was overlooked when he was trying to speak. I think it is important to remember that people with dementia can be anywhere within the hospital”
- “I have told all the departments that my husband has dementia and I noticed they became much more tolerant of his behaviour and mannerisms”
- “When I explained he had dementia it got a lot better, but I don’t know if I should keep having to tell them as it should be on his records”
- “The care was second to none, their dementia awareness was fantastic, and the staff spoke to my husband with dignity and respect and understood what he had said, even though he mumbles a lot”
- “We saw a consultant who was absolutely fantastic and explained everything to us which is so important for people with dementia”
- “I asked the staff about the ‘All About me’ document but the nurses had never heard of it and said they may just be on the older people’s wards”
- “They need to continue to improve awareness of dementia among staff”
- “For wards to become dementia-friendly, there is a lot of training available and it is excellent but people are so busy. Wards need to be more aware of the enhanced support teams and they need to use them at the right time”.

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Support at home:

Not all participants received support at home due to it not being needed currently. The participants who did receive support at home had mixed experiences depending on the provider. Participants who received care from DCC spoke highly of the care provided and commented around the ‘high level of training’. However, it wasn’t always available.

Our engagement officers were told some people do not qualify for DCC support, as their needs are not regarded as sufficiently critical. They also stated that when needs become more acute, people do not ask for a re-assessment, which means that people can find themselves in a critical state.

Participants who used private homecare services felt the carers were not always well trained, especially in terms of personal care. There were also concerns about the continuity of carers and flexibility around the timing of visits.

Sample of comments:

- “My mum was really used to the DCC carers; she was used to the routine and all the staff knew her and treated her with dignity and respect”
- “All of a sudden we were told the care would stop and we would need to have private carers” (DCC)
- “DCC have said we cannot use their carers”
- “Homecare staff should have proper training on dementia and how to treat people correctly”
- “Carers should turn up on time as it impacts on my wife’s tablets and medication as I have to wait until they have been before she can have her food and this means she is taking her tablets at all different times of the day. If they came on time there could be a proper routine”
- “The service is not reliable, there is no continuity and PLWD need to have the same staff so they feel comfortable and relaxed and do not get distressed”
- “I would like to have the same few carers so my wife will get to recognise them and she will be calmer and happier when she is being changed”
- “With the consistency of care being so important for PLWD there needs to be a move towards upping the status of a caring role by providing opportunities to build a successful, well paid career rather than the revolving door job it currently appears to be. Dementia care is a specialised area. If you get it wrong, the person being cared for suffers and their quality of life diminishes. With the right sort of care, life can be so much better”.

Link with occupational therapists and home adaptions:

Participants valued support offered by the OTs.

Some comments suggest waits for equipment to be installed following the assessment. We were told by a professional that there can often be delays in providing ‘dementia-friendly’ equipment such as coloured hand rails. The standard white equipment is readily available but orders have to be approved for customised equipment causing delays, therefore some people will just ask for standard equipment aids to help them at home.
Sample of comments:

- “An OT came to visit us. They did a very good job and we had a stair lift installed”
- “We had a number of people come to our home to see what aids and adaptions were needed. There was a lot of proactivity but we lost sight of it because of our daily battles”
- “We asked for handrails to be fitted to help dad move around the house, yet we are still waiting for them”
- “We asked for handrails three weeks ago and we are still waiting”.

Falls prevention:

We asked participants questions around reducing their risk of a fall. Their responses are shown in Fig. 1 below.

Participants who received information around reducing their risk of falls received the information predominantly through DCC, an OT or through the Living Well Programme.

Some participants were still waiting for their falls assessment.

Falls classes were found to be beneficial, but not everyone could attend due to difficulty with travelling to them.

Sample of comments:

- “It took a while. We contacted an OT for an assessment and five or six weeks had passed. We had not heard anything, so I chased them up and we were told we were 56th on the list”
- “Once I started them (falls classes) I have not looked back. They teach you how to get back up properly if you ever fall”.
Fig. 1:

Reducing the risk of falls

Safe and Well Checks (Derbyshire FRS):

Participants valued the advice and information received from the Derbyshire FRS, via safe and well checks.

Sample of comments:

- “The visit from the Fire Service was helpful as I had new fire alarms fitted”
- “They put new fire alarms in our house. This was nice to have and it was reassuring”
- “The Fire Service have been excellent and have supported us with a number of our patients” (Professional).

Support from day centres and independent groups:

Participants valued support offered by day centres and independent groups, such as church groups. Many participants explained day centres were a good place for their loved ones to go and participate in different activities and carers to receive respite.

A small number of participants commented around the fact that many groups and services provide support on the same days which in some cases causes people to have to choose.

A professional told EOs that workers are struggling to find activities for PLWD, and this has got worse in the last year.

Transport can also be a problem when it is not provided.

Sample of comments:
- “Day centres for PLWD are great. It is great for them to stay involved in activities but it also gives the carers a break”
- “We have two things this week but then we have nothing for the rest of the month”
- Jubilee Centre in New Mills: “It is excellent; all the staff are supportive of both my husband and I. They have provided me with great advice and have been helpful throughout our journey”
- Still Waters in Buxton: “We love it and there is so much community spirit. We get support from each other and the atmosphere is great”
- Age UK: “Age UK are excellent” and “the day centre is excellent. We like doing all the activities - we like to dance, sing, cook and play games. The staff are lovely, we are always doing something different”
- Midway: “It is a homely environment and doesn’t feel like a hospital. They are wonderful and they make it as nice as they can for you”
- Stepping Stones: “I cannot speak highly enough of them. They are so supportive and they have a person-centred approach. We used to visit and it would be £10 which included a three-course meal and it was available 9:30-3pmish. Transport used to be arranged for us and we would go on days out. We once went to Scotland. I didn’t have to do anything whilst we were in Scotland; they did everything for us and if it all got too much for me they were there for me”
- Stepping Stones: Following a discussion with a professional, she said that “there is a gap in service as the Midway hospital now only runs a two-hour day service. It used to be from 9-3. However, people are now dropping out as it is no longer viable for carers as it does not give them sufficient respite. There is great need for day care as it ensures that full-time carers get at least some time to themselves once a week. The end result may be that more people end up going in to a crisis and people have to be put in care”.

Derbyshire Dementia Support Service (DDSS):

Transition to Making Space:

Many participants had been involved for many years with its predecessors.

Most participants commented around the transition to the new provider. Some felt that they had not had a say in the change of provider and felt the transition caused a lot of distress as it was not done so well.

There was a lack of continuity with the DAs and there were ever-changing timetables and a general lack of organisation.

Sample of comments:

- “My mum has not coped well with the change and no longer engages at all”
- “My mum felt so uncomfortable and has asked that we do not go again”
“When they first took over it was not good but [named worker] is superb. I do not feel they get the backup they need. For example, the training, they have just been placed in the deep end”.

Training and knowledge of the dementia advisors (DAs):

A large proportion of participants felt the current DAs did not hold enough knowledge around dementia. In some instances, the DAs were not referring people onto other means of support such as Derbyshire Carers Association (DCA), Sight Support, or Strictly No Falling.

Sample of comments:

- “They (DAs) lack the insight. They say they have had dementia training but I feel they have been thrown into the deep end with very little support”
- “[Named worker (DA)] tries their best but he does not really know what he is doing and I would not feel comfortable contacting him for support”
- “When the Herbert Protocol came out we asked the facilitator if she knew about it. She explained she had been to the inauguration but did not have any information or forms for us”
- “Staff should be trained so they are able to offer dementia advice”
- “[Named professional] cannot seem to answer specific problems we are having around my husband’s condition and we are just told to go to the GP for help”
- “It was so disorganised and the workers had little knowledge of working with people who have dementia”.

Facilitation and quality of the groups:

The majority of participants felt that there was poor facilitation around activity sessions and support groups. Some activities such as bingo and quizzes were felt to be causing people to become upset and frustrated.

Participants explained they would prefer the groups to be a place of sharing information and learning about the support available, rather than social activities.

Some participants did speak positively of the groups and the experiences of the musical memory groups were mixed.

Sample of comments:

- “We told them we would like to have a focus but they just wing it”
- “The group facilitator seemed ill-prepared for two hours with people living with dementia”
- “My husband no longer wants to go. He feels the sessions underline what he can no longer do”
- “Stop just having colouring; I am not a child”
- “Why have things from before I was born? Of course I will not remember them”
“It is our first time at the group, we were not introduced and it was not made clear what the sessions were about. We were told it would be a place to get help but instead it was just playing games that we didn’t understand nor did we finish”

People do not all wish to be seen as the same. Some are happy and content to play bingo, and other games or discuss the 1940s. But others are highly professional retirees who have led lives that have never engaged in bingo. Maybe for this reason, some of the participants considered the sessions patronising and demeaning

“We want more than just games as we want to learn things that may help us”

“They should be more proactive and provide meaningful activities for people to attend and get something from it”

“More of a social rather than a source of direct meaningful support”

“They are excellent (groups). It is a lovely atmosphere and it is very relaxed with no strict agenda”

“My wife likes going to the group because it is for her and not her condition. I also feel very welcome when I go along too”

“It was a disaster; the worker who was tasked with taking the group would turn up late, leaving everyone without anything to do or any drinks. This was not just one occasion, this was every time. Eventually, the church who hosted the room offered to take over and facilitate it. It is now absolutely excellent with a turnout of around 40-50 regular members” (Musical Memories)

“They are really good” (Musical Memories)

“It was a smooth transition. I think it helped because it was the same people running the group. We both enjoy attending and it is an enjoyable experience”

“Making Space would not pay for a professional musician at a Musical Memories group. The song book they developed was too heavy for people to hold and had a large number of spelling mistakes in. There are a number of groups that have broken away and have started up on their own with support from local churches” (Musical Memories/Professional).

Confusion caused by changes to timetables:

There seemed to be a lot of confusion caused with changes to dates and venues with some being inaccessible. Also, some of the groups were changed from fortnightly to monthly.

Sample of comments:

“There are more benefits to regular sessions; monthly sessions are just putting people back into social isolation”

“The change of timings for the group is causing people to be unable to attend”

“The changes in times and venues has been an ongoing issue for the past year with ever-changing timetables and umpteen versions being printed and distributed. This is incredibly confusing for members, especially when some cannot always attend every month and find the times have changed the next time they turn up. They then feel unwelcome and that they have disrupted a group if the group has already started”.

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One-to-one support:

We were only able to gather one experience of a participant receiving one-to-one support. Participants in the main had received no meaningful one-to-one support, with quite a few participants explaining they had been ‘promised’ support which never happened.

Participants felt that they would have benefitted from one-to-one support, in terms of having a named person to go to for support but also, as a way of being effectively signposted to other services.

Sample of comments:

- “The staff do not know enough about dementia to provide accurate help. I have had to find out about things through friends and neighbours”
- “There should be a one-stop shop rather than ringing around different organisations and remembering who to chase up all the time”
- “When I was diagnosed I was referred to Making Space and they have been superb. They have been very helpful in providing support when we have needed it”
- “Initially they were wonderful. They told us ‘we can do this and help you with that’ and we felt absolutely supported and then we never heard anything from them. I contacted them and we managed to get a home visit. Again we were promised everything and received nothing. They are a waste of space. We have had nothing but false promises and I have no confidence in Making Space at all”
- “My wife is newly diagnosed and we have said in the group that it would be really nice for someone to give us a phone call, but the workers just seem to talk to other staff during the group. They do not come around to us and ask how we are doing”
- “Where are the resources to train the staff in providing one-to-one support in key issues which makes all the difference between crisis and providing the safety net through the tough times?” (Professional)
- “There should be a triage system for when someone is diagnosed where they can contact and make referrals on your behalf based on your needs so it is personalised. Alternatively, the triage system could offer signposting as sometimes you do not know who to turn to for support due to there being so many organisations involved”.

7.4: Living well:

Recognising the role of a carer and having an assessment:

Many of the participants spoke about how they now identified themselves as a carer for their loved one, but not all participants did.

There were mixed reports from carers with regards to access to information following their loved one’s diagnosis.
Those who did identify themselves as a carer were asked if they had received a carer’s assessment through DCA. This information is provided in Fig. 2 below which demonstrates a mixed response.

Some participants didn’t feel an assessment was needed or they could not see a benefit.

Some comments suggested a lack of communication with DCA.

Fig 2:

CARER’S ASSESSMENT

- Yes I have had an assessment
- No I have not had an assessment
- I am unsure whether or not I have had an assessment

Sample of comments:

- “My son explained to me how my role as a wife had now changed”
- “Yes, I am her carer, but she is my mum at the end of the day. She looked after me for 20 years so I am just returning the favour”
- “Yes, and I do everything for her and I want to carry on doing it for as long as I can. It is my job as her husband to look after her”
- “We are a couple and we support each other; that is what you do”
- “I had one (assessment) a while ago but I wanted to update it. I rang DCA to explain and was told someone would be in touch. A few weeks ago I contacted them again as I had not heard anything and I was told I was on their system but I have still not heard anything”
- “In the autumn of 2016 I had an assessment, however this was not finished and it did not lead to any further help. I had to do all the calling and arranging myself”
- “When I was waiting for the assessment to be done over the phone, I arranged for a befriender to sit with my husband whilst the assessment was taking place, but then DCA did not ring me back. It was very inconvenient as I had to arrange and change the time of when the befriender usually comes”
- “I can’t even get through to them. I have tried to call them a number of times in the last year and no one has rung me back. Carers don’t get called back and it is very vague as to what is on offer for them” (Professional).

Registering as a carer with a GP:
Most participants had registered as a carer with their GP, however other participants were unsure as to whether they had officially registered as a carer but believed their surgery or named doctor was aware of their caring responsibilities.

Some participants who had registered as a carer spoke positively of the support received by their GP surgery, using words such as “flexible” and “supportive.”

Sample of comments:

- “My GP is very sympathetic to my role as a carer”
- “The surgery have been very accommodating when I have needed appointments which fit around my role as a carer”
- “We have a carers champion in the practice and she makes sure that people have had a carer’s assessment and that they are aware of their rights” (Professional).

Carer’s emergency card:

With regards to the carer’s emergency card, responses were mixed with some participants explaining that they had a card but hadn’t had to use it, and some said they were not aware of the card, but would have found it useful.

Sample of comments:

- “I do not think there is enough emphasis put on emergency plans and not everyone has support around them”
- “This would have been useful. I was recently in A&E and I had no one to look after my husband so he had to come with me and sit there for seven hours. The staff were lovely but he was becoming very distressed”
- “I do feel alone at times and I worry about what would happen to my husband in an emergency”.

Respite care:

Participants explained they were aware of how to access respite but not all needed it currently. Some described the “lengthy” and “complicated” process to access respite, and the need for carers to be very proactive. Once the support was in place the experiences were generally positive, but not all carers felt they received enough respite.

Sample of comments:

- “I asked the care co-ordinator about respite and I was told about care homes. I didn’t want to put my husband in a home. I then found out after I had done some ringing around about a day centre and now my husband goes every Friday and he loves it”
- “I receive one week of respite per year, but apart from that I do not feel I receive enough support”
- “The only respite I have been offered is two weeks in a nursing home. All I want is an hour each week to get my head around things and this would mean I could carry on doing the things I need to do and not get overwhelmed”
- “I was unsuccessful in finding a befriender for my husband so I could no longer carry on with my art group and choir”.

Caring responsibilities having an impact on participants being able to care for themselves:

A number of participants felt that their caring responsibilities were having an impact on their own health due to difficulties accessing the relevant support. Others talked about feeling “unsupported” and “alone” when caring for a loved one.

A number of carers felt they had lost their identity due to their caring role and were no longer able to participate in activities they used to enjoy.

A number of participants from a group in South Derbyshire felt that it was important that carers received a free flu jab, explaining that “if we became ill it would have a big impact on the health and social care system so we should be allowed free flu jabs. Some surgeries in the area are very proactive and encourage carers of people with dementia to have a flu jab but others do not”.

Sample of comments:

- “My GP advised me to access support from a CPN for my husband whilst I was going through treatment. The CPN got in touch two weeks before my treatment was finished so for me it was too late, the moment had passed. It is no one’s fault, it is just the system was too slow”
- “Unwillingly I am too proud to admit when I need help for my wife and I get frustrated as sometimes I have difficulties going for my own appointments as they do not fit around my caring responsibilities”
- “They keep telling carers to look after themselves but you just can’t”.

Support for carers:

Many carers felt they just needed a little more help to get the right support in place, i.e. in terms of time and understanding. Carers felt they had to become proactive in order to get support.

Sample of comments:

- “Yes, I had a lot of support from DCA; they are very helpful in supplying literature about carers groups and what is in the area to help”
- “I have had to find out everything for myself”
- “I had to sort it all out myself. I was given brochures which took a long time to go through. I eventually found someone who then came to our home three days a week; they were lovely and really helpful.”
- “Whilst you cope you are left alone”
“Family carers need emotional support. It is devastating walking alongside someone as they lose their independence and become increasingly reliant upon their carers for every aspect of living during every step of their distressingly slow demise”

“It really takes a toll on me. It is emotionally and physically draining; no one really understands the extent we have to go through to make sure mum is OK”

“There are more and more people living at home with dementia and as a carer I feel I was taken advantage of. It isn’t easy making phone calls and keeping in touch with services when you are caring for someone”

“We are still people too, but all I am now is the person who looks after my husband 24/7. I am not me anymore; I do not count”

“Let us understand the decision making. Give us the time we need and the support. Our worlds are being turned upside down. If we get upset or angry please forgive us and understand why we are feeling like we do. We are grieving now and then we will have to do it all over again when our loved one passes away”.

Co-ordination of services to support carers:

Support for carers was described as disjointed. Carers asked for better co-ordinated services, and better communication, i.e. a central organisation that can be contacted for advice, signposting and referrals.

Many participants who were caring for a loved one felt that they had to be very proactive and become ‘experts’ to be able to carry out their caring responsibilities and often felt this was ‘overwhelming’ and ‘confusing’ due to receiving very little support from organisations.

Carers also felt it should be the role of professionals to keep in touch, and check out how they were doing, rather than the carers doing all the chasing. Participants felt there is a need for more peer support.

Sample of comments:

- “Everything (services) seems disjointed and you are unsure of who to go to”
- “There should be a central organisation who you could contact for advice and referrals if required”
- “There should be a triage team where you can contact them for advice, signposting and referrals”
- “There are too many organisations which makes enquiries for help, lengthy and complicated so people are often left to fend for themselves”
- “They rely on people being forceful but not everyone is like this and people are struggling with everything. I don’t like the fact you have to ask for every bit of information or help when it should be offered”
- “My caring duties leave me little time or energy for organising and searching things out and during my occasional free time I need that to relax and recuperate”
- “Organisations should be more proactive (i.e. making contact with carers, calling to see how they are) which would take a bit of pressure away from carers”
“I would like to have someone to talk to as I feel alone. I have to carry on for my husband as there is no one else to care for him. I would like to be able to talk to people who are going through the same situation on a one-to-one basis because professionals never have enough time to talk to us”

“Peer support would have helped, being able to talk to someone who is going through the same as you and knows what to do. It would be good to set up a buddy system so that things are shared and you can have someone to talk to”.

Paying for care (self-funding):

Some participants who were self-funding explained their struggles of getting the advice on relevant care and support. Participants explained they had contacted the relevant agencies for support but as they were self-funded they were often just sent leaflets or brochures through the post with no further support.

Sample of comments:

- “Just because we have money does not mean we know how to care for someone with dementia properly”
- “The moment they found out we had money, our case was closed within a week”
- “Left to fend for ourselves”
- “I needed to go for an appointment for my own health and the prices I was getting back were so expensive. The figures were ridiculous for a few hours. The appointment was for my own health; I was not going on a jolly”
- “There should be more support for self-funders as it is chaos and very frustrating”
- “I am happy to pay for day care but people on the phone do not know where to refer you. I have spent hours on the internet looking at what help is available to us”
- “If you have your own funding, you are on your own. If my husband had cancer he would have been treated under the NHS but with dementia they know there is no hope of you getting better”
- “It seems to me that because we were self-funding we were just left to get on with it, but with dementia there can be a rapid decline which can mean we need help and we need it now but I don’t feel the system accounts for that”
- “It felt as though your mum has a carer, me, and you have money so you can manage on your own”.

Paying for care (Continuing Healthcare):

A number of participants expressed their concerns around understanding the CHC process, and difficulties accessing funding. There were also comments about jargon used at ‘funding meetings’ which made them feel unable to participate.

Sample of comments:

- “It has taken me over four months to try and understand the CHC process. Professionals have made no effort to help me understand the process or to get funding that we are entitled to”
- "It has been an absolute nightmare; it is a confusing process due to the lack of information and it is very easy to be fobbed off, so if you are not aware of the process you can easily miss out"
- In late 2017 one participant’s husband was discharged from hospital on End of Life Care (EOL) and said that “they did not fast track him as they can only do this if they suspect EOL within six weeks so I had to pay the care costs and I did not get CHC until two days before he passed away”
- “The funding meetings are so important and cause so much stress to families. They should provide people with more information prior to the meetings so people are aware of how the process works and what to expect”
- “We are a strong family; we will challenge the NHS or Social Care but not everyone will. What would have happened to them?”
- “We were told we would be eligible for £530 per week but her care was around the £1000 mark. That would mean as a family we would need to fund £400 per week. How could we afford to continue to fund that, when she is still quite young? It was a very challenging time for the family”
- “We tried to challenge the decision but the social worker kept on saying ‘you can get her nursing care for that price’. Yes, maybe we could, but we did not feel they were fit for purpose; she needed dementia and nursing care”

➢ Dementia Friendly Communities:

Aware of dementia-friendly communities:

We asked participants whether they were aware of Dementia Friendly Communities (DCF) and whether they thought Derbyshire was dementia-friendly. Some participants gave some suggestions for improvements.

The chart in Fig. 3 shows the responses for the people who answered this question.

Fig. 3:

Aware of Dementia Friendly Communities:

Sample of comments:

- “It is getting better”
- “They are doing what they can”
- “For people to have more of an understanding and to be patient; I do not want to shout ‘this man has dementia’, and when we go out for meals it can take him a long time to choose his food”
- “When we get a diagnosis it doesn’t change who we are. We don’t want to be treated any differently. We don’t want people to start walking off the minute dementia is mentioned”
- “I would like shops to count out change. My husband gets very confused as he is just given a handful of change”
- “To have a leaflet on what dementia-friendly means. This way people would be more aware and we would be able to spot things”
- “To have dementia-friendly parking spaces; my husband does not qualify for a parking badge and I have to make sure I park somewhere where I can open the door wide enough for him to get out”
- “A friend has been to the airport and if you are with someone with dementia they give you a lanyard which makes people aware they have memory loss. I think this would be good if shopping centres and supermarkets could do something like this”
- “For supermarkets to have a dementia-friendly shopping session. I have heard they do this somewhere in the north of the county”.

Involvement in dementia-friendly projects and groups:

A small number of participants were involved with dementia-friendly projects and participants who attended dementia-friendly swimming or walking spoke very positively of them.

Sample of comments:

- “It is brilliant; it enables the person with dementia to really open up about how they feel” (Amber Valley Discussion Group)
- “I love the swimming. I think the more I can do, the better I will stay” (Dementia Friendly Swimming)
- “The dementia-friendly swimming has been a boom to my husband. When he was first diagnosed I tried for ages to get him back in the pool. He was a beautiful swimmer but he just would not swim for me, then I heard of this and when he got back in that pool he took off. People here are lovely and it is great for us to meet new people and talk to other carers”
- “I go on the dementia walks which are good. You can just walk and talk together and it is really nice”.

Experiences of People Living with Dementia under the age of 65

7.5 Preventing well:

Education and consideration to images used:
As for the over 65s, many participants explained there needs to be a shift from the focus around Alzheimer’s disease to awareness of all types of dementia.

Participants under 65 years old also felt the image used for dementia often portrays a “negative image” and can sometimes imply that dementia only affects older people.

Sample of comments:

- “They always use images of older people who cannot eat or cannot do something for themselves. This is just portraying a negative image”
- “It would be better if there was a wider range of materials available so people are aware of other types of dementia and the symptoms to look out for”
- “If you are young you cannot get it recognised as dementia. My husband did not want to recognise it himself because when the word ‘dementia’ is used you automatically think of an older person, and they need to change their image”

7.6 Diagnosing well:

Initial discussion with a professional:

When participants discussed their initial concerns with a GP, the experiences were mixed. Some participants felt their symptoms had been ‘downplayed’ and were told they are “too young to have dementia”, whereas other participants had very positive experiences and felt listened to by their GP.

Sample of comments:

- “The doctor was very sympathetic and understanding towards me and made a referral”
- “They were really good. They spent the time listening to me, they completed a questionnaire and then referred me to the MAS clinic”
- “They downplayed my symptoms and said that I will be fine in a week or so”
- “Not really, they kept on saying I was too young and then they put me on tablets for depression for around a year”
- “I wish I didn’t have to keep on going back to my doctor before I got a referral just because I was too young”
- “No, we feel we have been left in a corner as my husband is younger. Lots of professionals have been in touch to say they will help but most of them we never heard from again”.

Information following diagnosis:

When asked if they had received adequate information following their diagnosis, participants explained the difficulty in accessing information around young-onset dementia and often did not receive the information required. Whereas, with the over 65s, some participants received ‘too much’ information.

Sample of comments:
- “It is so difficult to get information around early onset”
- “After diagnosis you are told about different individuals that will be involved in your dementia journey but there are too many to remember. It would be good if once you are diagnosed, you received a one-page ‘road map’ that shows who is who and their role in your journey”

7.7 Supporting well:

Support for young-onset:

Many participants felt unsupported and felt this was due to their age as there seemed to be a lack of support and activity sessions for younger people.

Participants explained the importance of recognising young-onset dementia and for the system to be able to provide meaningful and tailored support.

There was a discussion around the importance of ‘understanding what dementia means’. As one participant said, “people do not understand what dementia means, there is no support available for people because it is an invisible disability”.

All the members praised the DEEP groups and explained it allows them to be themselves and be with people who have some understanding.

Sample of comments:

- “No, I feel we have had little support. After the Living Well programme nothing else was offered and the only support available is where people are of a much older age, for example, luncheon clubs. Workers seemed to give up as there was nothing for my age”
- “I am not something you put into a box - I am a person”
- “It has taken all my confidence away and I do not want to go out”
- “We tried a lunch club, but this is for very old and ill people and my husband felt very uncomfortable as he saw what he may become”
- “Not everyone wants a diagnosis but without the diagnosis there is no support”
- “Nothing is geared up for people my age. Nobody will talk to us or give us information and it makes me feel invisible”
- “I feel I am banging my head against a brick wall because the GP does not recognise what I am saying”
- “I feel like services have treated me like an idiot because I am young”
- “I feel people start talking to my mum when they find out I have dementia, just because I am young they don’t know what to say to me”
- “It is like a family. [Named professional] helps me so much I am really grateful for the support” (DEEP Group).

Improvement for people with early-onset dementia:

Following a number of discussions with participants with young-onset dementia, participants made some suggestions for improvements they would like to see in the future.
Summary of comments:

- **More support groups for younger people with dementia**
- **Improve nurses’ attitudes, let people know that younger people can have dementia as well.** One participant claimed that there was no dignity or respect for her husband as they just thought he was being awkward
- **Personal health budgets to do age-appropriate things rather than quizzes and bingo**
- **Support for families of people with young-onset dementia, especially their children**
- **Increase employer’s knowledge of accommodating someone with young-onset dementia in work**
- **Financial support, as often people with young-onset dementia are not entitled to a state pension, and have not saved enough in private pensions.**

**Experiences of people living with dementia from the BME community:**

**7.6 Diagnosing well:**

**Receiving diagnosis:**

Participants talked about the difficulty getting a diagnosis of dementia, taking up to two years for the formal diagnosis.

It was explained that some individuals are able to get a quicker diagnosis when family members can engage with services, however sometimes families can be in denial.

Sample of comments:

- “People often resort back to their mother tongue when they have dementia and although interpreters are available, there can be a ‘lack of trust’ along with ‘cultural differences’ with interpreters which prevents access”.

**7.7: Supporting well:**

**Support for participants from the BME community:**

Participants felt there is also a “huge language barrier” with BME carers and PLWD in hospital settings. Participants felt it was important to have effective communication with PLWD as they can be often ‘in denial’ and will be ‘complacent’ with their answers.

It was discussed that homecare workers seem to lack general dementia awareness, which is also evident in numerous sections of the report.
It was discussed that individuals from the BME community need to have tailored support to meet their needs and one participant suggested having more befrienders from their local community.

Sample of comments:

- “One of our members was in hospital and there was a huge language barrier and coupled with dementia it made it a very isolating place. The individual was then discharged late at night, returning to an empty house which caused more confusion”
- “We feel they need more training in communicating with people living with dementia. They ask questions such as ‘do you want a bath?’ to which they respond ‘no, I am OK’. This happens with a number of professionals; they do not probe enough”
- Participants also explained they often felt care was ‘rushed’ as they were only provided with 10-15 minutes of care, which was not enough time to support individuals effectively.

8. What should happen now?

1. Continue to work to improve the provision of early education to raise awareness and understanding of dementia
2. Increase information and communication around all types of dementia and provide clear information on all signs and symptoms
3. Consider the images used for literature and campaigns to ensure the message is clear that dementia can affect anyone, regardless of age
4. Ensure initial concerns around suspected dementia are listened to:
   - Regardless of age
   - Consider the concerns expressed by carers and family members.
5. Work to reduce language barriers to ensure this does not delay diagnosis and to ensure people are fully supported throughout their journey
6. Consider and evaluate the amount of information provided following diagnosis, and to consider whether the information is delivered at the right time
7. Ensure there is opportunity for PLWD, carers and family members to ask questions following the diagnosis:
   - At a later date, rather than immediately after diagnosis
   - Ensure carers and family members are able to ask questions around their loved one’s diagnosis in private.
8. Work to ensure all PLWD, their carers and family members are signposted to appropriate support following diagnosis and ensure the support is easily available to access at a later date, if not required immediately after diagnosis
9. Promote the Living Well Programme and the question and answer sessions
10. Ensure the care and support needs of PLWD are met and regularly reviewed, as suggested in the Dementia Well Pathway
11. Ensure workers and professionals are able to identify patients/clients with dementia and their carers/family members and have received adequate training to be able to meet their care and support needs in all settings
12. Maximise the need and importance of continuity, routine and training for homecare staff
13. Improve the experience of people involved with the DDSS in terms of more appropriate activities, more training for staff and ensure the one-to-one support is readily available
14. Develop a role of a ‘named person’ to enable people to have one point of contact to be able to access the advice and support needed
15. Work to address the need for more support and activities for PLWD under the age of 65
16. To ensure carers receive the correct level of support to effectively care for their loved one and also look after their own health and well-being
17. Improve co-ordination of support services for carers, so they know where to go for advice, signposting and referrals
18. Ensure families who are self-funded have the correct level of information and support
19. Address the lengthy process for accessing respite care and to make it easier to access
20. Provide clear information around Continuing Healthcare and the use of jargon, so people are able to easily understand the process.
9. Response from service provider

The below table provides a summary of the responses received by service providers and commissioner’s for the Healthwatch Derbyshire Dementia Report. The table has been produced due to the volume of responses received.

If you would like to view a full version of the responses please visit: https://healthwatchderbyshire.co.uk/wp-content/uploads/2018/05/HWD-DEMENTIA-REPORT-FULL-SERVICE-PROVIDER-RESPONSES.pdf

Alternatively, you can request a hard copy by contacting us:

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Sustainable Transformation Partnership (STP):

NHS organisations and local councils have come together to develop plans for the future of health and social care. Derbyshire’s STP, Joined Up Care Derbyshire, brings together work that has been taking place across the county to coordinate services better so they support people to stay well. For more information on Derbyshire’s STP please visit: https://joinedupcarederbyshire.co.uk/

What should happen now?

<table>
<thead>
<tr>
<th>Response’s:</th>
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<tr>
<td><strong>1) Continue to work to improve the provision of early education to raise awareness and understanding of dementia.</strong></td>
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**Sustainable Transformation Partnership - Dementia & Delirium Workstream:**

The STP has a key goal to minimise the risk of people developing dementia and delirium in dementia, and improving pre-diagnosis information and health promotion for prevention of dementia are important steps to this goal. Partners within the STP across the health and social care, and voluntary sector, have a stated commitment to improving the availability and quality of information for early education, awareness raising and the various types and manifestations of dementia. Partners also have a stated ambition to ensure their work is integrated to further support this, and other key STP goals. The STP, through the framework of operation, will monitor how this goal is
progressed and achieved (This action also relate to recommendation 2 and 3).

**Derbyshire Clinical Commissioning Groups (CCGs):**

- Specific sessions relating to ‘early recognition and early intervention of dementia’ were commissioned in Erewash in November 2017 (This also relates to recommendation 2)
- Dementia Friends training is continuing across the county and many sessions are being offered as part of the Dementia Action Week (May 21st - 25th)
- The CCGs are part of the local and county Dementia Action Alliances (DAAs) and are working on plans for dementia-friendly communities (DFC). Members of these alliances include retailers, banks, pharmacies, local MPs, and Councilor’s and local businesses.

**Derbyshire County Council (DCC):**

One of the key commitments outlined in the DCC Council Plan is to develop support services for people living with dementia (PLWD) to help them live well and independently. DCC will continue to provide opportunities for its staff to attend awareness raising sessions to improve understanding of dementia.

**Chesterfield Royal Hospital NHS Foundation Trust (CRH):**

CRH aims to have an outstanding ‘dementia-friendly’ culture of care which will be achieved by ensuring that all Trust staff are dementia aware. The Trust will continue to run and promote an extensive portfolio of dementia training, which currently includes:

- Induction - all staff receive dementia awareness training at induction (also relates to recommendation 11)
- Annual essential training - dementia awareness training for all staff (also relates to recommendation 11)
- Care Certificate training - a nationally recognised qualification, including an agreed set of standards setting out the knowledge, skills and behaviours expected of Healthcare Assistants (HCAs)
- Promoting Quality in Dementia - provides a better understanding of dementia and lived experience; this helps to equip staff in providing appropriate care and supporting people to live well with dementia - available for all staff
- Best Practice in Dementia Care - in line with national dementia strategies, this programme helps organisations to achieve national and regional care standards. The
programme is accredited by the Royal College of Nursing with 16 CPD hours and recognised by Skills for Care.

**Derby Teaching Hospitals NHS Foundation Trust (DTHFT):**

The Trust’s dementia awareness training is now mandatory. Therefore, all staff starting in the Trust will receive the training as part of their induction. The training is also available to any existing staff who have not done the training since it went mandatory, via the one stop shops. The Lead Nurse for Dementia is also able to provide training to any areas where required.

**Derbyshire Community Health Services NHS Foundation Trust (DCHS):**

DCHS is developing a Dementia Strategy which will include the following key strategic objective: the continued development of our Trust as a dementia-friendly organisation, with environments that promote better outcomes, and that are safe.

| 2) Increase information and communication around all types of dementia and provide clear information on all signs and symptoms | **Derbyshire CCGs:**  
Dementia-friendly practices exist in some areas across the county (7 out of 12 in Erewash and 3 in the north). This continues to be rolled out but is a slow process.  

**DCC:**  
DCC will continue to improve and develop a range of information and signposting to improve pre-diagnosis information and health promotion for the prevention of dementia.  

**Derbyshire Healthcare NHS Foundation Trust (DHCFT):**  
At the point of diagnosis all patients are:  
- Informed of their specific diagnosis and the rational for why this is an appropriate diagnosis  
- Offered either face to face or referral to Making Space the Derbyshire Dementia Support Service (DDSS). They are commissioned to provide literature including information specific to their diagnosis  
- Offered the Living Well with Dementia education course that provides further education on types of dementia.

DHCFT will continue to:
• Work alongside DDSS staff in the diagnosis clinic
• Provide information and access to the Living Well with Dementia education sessions
• Promote Dementia Q and A sessions in different formats.

CRH:

In addition to the above training (in recommendation 1), CRH has several avenues of support, where staff, patients and carers are able to seek further information:

• The Trust has established an Enhanced Nursing Support Team (ENST) to offer a gold standard of care. The ENST is a dedicated team of HCAs that provide a level of increased supervision to patients, including those with severe dementia. The team have received specific training and reside in a virtual ward managed by the Older People’s Matron. This provides a flexible workforce with staff allocated on a shift by shift basis
• A Trust Older Person’s Team is available for support, advice and assessment
• The Trust also has dementia information resources and packs available from Library Services
• Dementia drop-in sessions are held every Tuesday afternoon and are currently run by Making Space - patients and carers are able to access information, advice and support about dementia
• In January 2018, a Carers Liaison Officer came into post, who offers support, information and signposting to carers, and works with staff to ensure that they are able to provide this support
• The Trust also has support from DHCFT Liaison Team.

DTHFT:

The Lead Nurse for Dementia is currently organising a study day for staff in the Trust, around the different types of dementia, which will include looking at the signs and symptoms. This will help staff to feel confident to share the information with other staff members and carers. Staff can also signpost relatives, patients and carers to the information hub where there is information available around dementia. Information on types of dementia and signs and symptoms is to be provided via the dementia portal for all staff on the intranet. Enquiries are to be made to see if information around types of dementia and signs and symptoms can be put on the Trust web page for the public to view.
### Derbyshire CCGs:

We are working closely with Making Space and Alzheimer’s UK who are supporting us with the production of literature for activities and events such as those taking place for Dementia Action Week in May. The current service provider, Making Space, has reviewed and reproduced its literature in light of these recommendations.

### DCC:

- DCC Public Health commission the Alzheimer’s Society to deliver the DFC project, highlighted in the report, that uses stakeholder engagement and awareness raising to empower people, businesses and organisations to become more dementia-friendly and help PLWD maintain their independence in the community.
- DCC has expanded its ‘Safe Places’ scheme to include those PLWD.
- The DDSS, provided by Making Space, plays a key role in disseminating information about all types of dementia, and DCC are working closely with Making Space to ensure the service reach is as wide as possible, including affected younger people.
- The Adult Care Information Service will review the leaflet and web page that it produced about dementia to take account of the point raised about the images used. We will also consider adding more information about younger onset dementia to our information set.
- The Carers in Derbyshire website ([www.carersinderbyshire.org.uk/](http://www.carersinderbyshire.org.uk/)), which has been developed by DCC, will be reviewed to ensure there is sufficient information provided about younger onset dementia.

3) Consider the images used for literature and campaigns to ensure the message is clear that dementia can affect anyone, regardless of age.
CRH:

A key part of Trust dementia training is to discuss the potential for patients under the age of 75 to have dementia, to equip staff with the knowledge and understanding that dementia can affect all ages. The Trust aims to be inclusive with all images used in publications including those regarding dementia. The Communications Team have been made aware of this recommendation and the Older Person’s Team will ensure they liaise with this team when producing any future publications for dementia at the Trust.

DTHFT:

The study day organised by the Trust will include looking at early onset dementia. We will also look at resources and information around early onset dementia, available to both staff and the public in the information hub.

| 4) Ensure initial concerns around suspected dementia are listened to: |
| - Regardless of age |
| - Consider the concerns expressed by carers and family members |

Derbyshire CCGs:

Awareness raising activities are taking place including:
- An individual with dementia and his wife providing a presentation at the Practice Managers Meeting in Erewash, to share their experiences in order to raise awareness
- Dementia patient and carer stories being heard at the Governing Bodies of the CCGs to raise awareness of the lived experience
- The findings of all the reports and other service concerns being fed back through services, for learning
- The Memory Assessment Service (MAS) has been newly configured and decommissioned to have one MAS for the county and a MAS 24 service. The MAS at Ilkeston hospital underwent a quality review visit at the end of last year and was evaluated very well with a recommendation to gather patient and experience feedback going forward.

DCC:

DCC adult care assessments are needs led and take into account any concerns regardless of age. DCC will continue to ensure that all services provided or commissioned by the Council are responsive to concerns expressed by carers.

DHCFT:

DHCFT provide training to GPs and the wider health community
as requested. DHCFT will continue to respond to requests to provide dementia training at Quest events and to the wider health community.

**CRH:**

The Trust takes initial concerns around suspected dementia seriously and are able to listen to patients and carers via several avenues, including:

- Part of the Trust’s Older Person’s Team role is to find, assess and refer patients with dementia. All patients over the age of 75 will be assessed within 72 hours of admission to hospital. Those under 75 are assessed on referral from staff or self-referral.
- The Trust is establishing Dementia Champions, with the aim that every ward and clinical environment in the Trust will have a nominated champion to act as a link to the Trust developments in dementia, provide support and training to staff and work as a champion for excellence in care.
- The Carers Liaison Officer is able to offer one-to-one support, advice and signposting to carers.
- The Trust has signed up to John’s Campaign, the principles of which recognise the valuable contribution of carers to the health and wellbeing of dementia patients.
- The Trust’s Carers’ Charter pledges that we will support the needs of carers, and work with them as partners in care. The Trust has developed a Carer’s Passport to support this.
- 24/7 support is available from DHCFT’s Liaison Team.

**DTHFT:**

Dementia link nurse/champions meetings are being re-established, so we have lots of champions across the Trust who can address any concerns raised by families and carers.

| 5) Work to reduce language barriers to ensure this does not delay diagnosis and to ensure people are fully supported throughout their journey | **STP Dementia & Delirium Workstream:**

The STP workstream is in the process of developing an engagement strategy, and reducing the impact of language barriers and improving response to the needs of our BME communities across Derby and Derbyshire is key to that development. Health and social care partners are considering integrated ways of working they could deliver to reduce language barriers and improve support for people living with dementia. Progress will be monitored through STP framework at strategic, implementation and monitoring levels. *(This action also relates to recommendation 4).*

V1 Dementia Report 22/02/2018 HM
### Derbyshire CCGs:

Provision of information in other languages is greater in the city and we will work across the county to look at this and assess need. With regards to communication generally, the dementia-friendly training covers general communication with people with suspected dementia and/or memory issues.

### DCC:

The Council are working with key partners, in particular the BME forum to consider how to respond more proactively to the needs of BME communities across Derbyshire. All services provided or commissioned by DCC are required to facilitate access for those who may experience language barriers.

### DHCFT:

There is no evidence to support the belief that BME patients are delayed in the process of diagnosis once referred to MAS. DHCFT provide Urdu and Punjabi dementia Q and A sessions and Living Well with Dementia Education Group. DHCFT will continue to promote the services in appropriate formats.

### CRH:

The Trust has access to interpreting services to support communication with patients and carers.

### DTHFT:

The Lead Dementia Nurse will ensure that staff are aware that interpreters can be booked for patients where there is a language barrier, when discussing a diagnosis of dementia and when attending appointments. We can also look at making sure that written information is available in various languages to reduce any language barriers.

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<thead>
<tr>
<th>6) Consider and evaluate the amount of information provided following diagnosis, and to consider whether the information is delivered at the right time</th>
<th><strong>STP Dementia &amp; Delirium Workstream:</strong></th>
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<tbody>
<tr>
<td>As the provision of information around preventing well develops, the volume of information useful at the point of diagnosis can be reviewed. As uptake of the Living Well Programme improves the volume of information provided at the point of diagnosis may also reduce. There is a stated commitment by health and social care partners to work in an integrated manner to deliver to the information needs of individuals, and to review the way in which this happens in an ongoing manner. The STP implementation group will provide a</td>
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format to enable review and monitoring of progress. *(This action also relates to recommendation 7).*

**Derbyshire CCGs:**

Feedback from PLWD in Erewash identified the need for a local directory to support them/their family post diagnosis *(This also relates to recommendation 8, 11 and 17).* We are also working with Derbyshire Carers around the general level of information given to carers at assessment and throughout the journey, especially as circumstances change. This is to improve the timeliness of information. There is also a specific intention to look at the information and support given when circumstances change and people need to consider care and residential support, as this is often a time of pressure to make decisions in a short time period. Making Space are aware of this and work with their advisors to look at how they support this stage of the journey.

**DCC:**

In partnership with the NHS and others, DCC will review the information made available following diagnosis, to ensure people do not feel overwhelmed. The review has commenced and already identified that the DCC ‘Guide to Dementia Services’ leaflet should be provided at all MAS clinics.

**DHCFT:**

Information leaflets are provided by DDSS and the information is agreed with Commissioners.

**CRH:**

The Trust provides patients and carers with information regarding dementia that they are able to take away and read at their convenience. There are also trained staff members available to offer additional support, information and signposting to patients and carers as appropriate.

**DTHFT:**

Considerations and discussions to be made around what information is provided to patients and families following diagnosis, and whether this information is appropriate at the time of diagnosis.

**Making Space/Derbyshire Dementia Support Service (DDSS):**
DDSS provide an option for support immediately following diagnosis at MAS across Derbyshire. Where information, guidance and support is requested at that time by a PLWD and/or carer, this is provided. In addition, the option is given for a one-to-one home visit from the same Dementia Advisor (DA). For anyone that wishes to receive support at a later date, the service information leaflet and contact details of the DA in attendance is provided. DDSS will review this process and establish whether a follow-up mechanism would be beneficial when the service is declined at this initial stage (this also relates to recommendation 7).

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<tr>
<th>7) Ensure there is opportunity for PLWD, carers and family members to ask questions following the diagnosis:</th>
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<td>- At a later date, rather than immediately after diagnosis</td>
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**Derbyshire CCGs:**

This forms part of the MAS/diagnosis pathway, where individuals are offered attendance at specific Living Well sessions. There is also an intention to look at the contents of the Living Well sessions, as some people have said they would like a refresher or an update later on in the journey, or as needs change. The MAS has developed so that nurses can give the diagnosis, although this diagnosis will have been made by the consultant following a review of the assessments. This allows for greater time for the family and the patient to discuss the diagnosis and there are options for them to come back and see the nurse for more discussions at a later stage. There is also the option for them to arrange to see the consultant at a later stage, and telephone discussion and conversation with the team is also encouraged. Dementia Q and A sessions are also held at Ilkeston Community Hospital.

**DCC:**

Staff from the DDSS attend all MAS clinics to offer information and advice straight after diagnosis, or arrange an appointment at a later date. Private appointments are also offered to carers/other family members. DCC are continuing to work closely with Making Space and DHCFT to ensure the offer of information/advice following diagnosis is timely and proportionate.

**DHCFT:**

Dementia Q and A sessions are available. The Living Well with Dementia Education Group is offered to anyone diagnosed with dementia and their carer/next of kin. DHCFT takes the clear view that the diagnosis belongs to the person diagnosed and it would be inappropriate to discuss a specific diagnosis without the person diagnosed being present.
<table>
<thead>
<tr>
<th>CRH:</th>
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<tbody>
<tr>
<td>The Carers Liaison Officer, Older Person’s Team, DHCFT Liaison Team, Carers’ Champions and Dementia Champions are available to answer questions and provide further information whilst patients/carers are at the hospital, including one-to-one discussions. Trust staff will also signpost to support services in the community.</td>
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<table>
<thead>
<tr>
<th>DTHFT:</th>
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<tbody>
<tr>
<td>We will look at what follow up appointments or resources are available to patients and families following diagnosis for any follow up questions after initial diagnosis. We need to ensure that patients and family members have an opportunity to ask questions independently and privately following diagnosis.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>8) Work to ensure all PLWD, their carers and family members are signposted to appropriate support following diagnosis and ensure the support is easily available to access at a later date if not required immediately after diagnosis</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>STP Dementia &amp; Delirium Workstream:</strong></td>
</tr>
<tr>
<td>Working to ensure all PLWD, their carers and family members are signposted to appropriate support following diagnosis and ensuring that support is easily available, is key to the activity of the STP workstream. Partners have a stated commitment to working in integrated ways to achieve this goal, not only within the bounds of what is available but also in seeking to advocate further for funding and services to support that aim. The STP strategy and implementation partnerships will be key to this activity.</td>
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<thead>
<tr>
<th>Derbyshire CCGs:</th>
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</thead>
<tbody>
<tr>
<td>The MAS has started to provide this level of signposting, with referral and signposting to the current service provider (Making Space).</td>
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<thead>
<tr>
<th>DCC:</th>
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<tbody>
<tr>
<td>As part of the dementia workstream of the Derbyshire STP, we will work with partners to ensure signposting and support for PLWD, their carers and family members is timely, appropriate and accessible.</td>
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<th>CRH:</th>
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<tbody>
<tr>
<td>The Carers Liaison Officer offers a follow-up phone call with carers two weeks following discharge of the PLWD to ensure their support needs have been picked up in the community. There is also a self-referral slip for Derbyshire Carers within Trust carer’s packs. Trust staff are trained to recognise and identify carers both at induction and annual essential training.</td>
</tr>
</tbody>
</table>
There are dementia and carer’s links provided on the Trust web pages, which signpost to further support.

**DTHFT:**

We can provide all patients and relatives with a resource folder which contains lots of information around dementia and also signpost them to the various community groups who they would be able to get support from.

**DCHS:**

DCHS is developing a Dementia Strategy which will include the following key strategic objectives:

- Early specialist support for people who have been diagnosed with dementia
- Targeted support (including cognitive stimulation and meaningful activity) to enable people with moderate dementia to continue to live well
- Listening, involving and engaging with PLWD and their carers including the timely completion of dementia care plans
- Ensuring prompt referrals to specialist services

<table>
<thead>
<tr>
<th>9) Promote the Living Well Programme and the Question and Answer Sessions</th>
<th><strong>STP Dementia &amp; Delirium Workstream:</strong></th>
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</thead>
<tbody>
<tr>
<td><strong>STP Dementia &amp; Delirium Workstream:</strong></td>
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<tr>
<td>The transformation of day hospital services is a key goal of the STP and within this there are changes planned to the way the Living Well group is offered, to improve take up. It is planned that individuals diagnosed with dementia will be contacted up to three times during the year following diagnosis to encourage take up of the programme. The STP key goal is to ensure that services available anywhere in Derby or Derbyshire are equitable and this transformation is planned for North and South services in partnership with the two Trusts who will deliver the service. It should be remembered that the Dementia Q and A sessions are run voluntarily by DHCFT staff, and although they work very hard to sustain the offer, it is not always available. Under the STP, partners have agreed to look at ways that this could become more sustainable as it is a valuable service.</td>
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<tr>
<td><strong>Derbyshire CCGs:</strong></td>
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<tr>
<td>This now forms part of the MAS/diagnosis pathway where individuals are offered attendance at specific Living Well sessions. We will work with DCC, DHCFT and DCHS to promote</td>
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</tbody>
</table>
the Living Well Programme and Q and A sessions more proactively.

**DCC:**

DCC will work with DHCFT and DCHS to promote the Living Well Programme and Question and Answer sessions more proactively. Staff from the DDSS, commissioned by DCC, already attend both Living Well, and the Q & A sessions, and we will consider options to integrate the service even more closely.

**DHCFT:**

DCHFT run the Dementia Q and A sessions in the South. They are due to recommence in the North (10/18). The Living Well with Dementia Group is provided by DHCFT in the South and is offered to all people newly diagnosed.

**CRH:**

The Trust:
- Will continue to work closely with DHCFT Liaison Team, who would refer into the Living Well Programme on behalf of the Trust, at diagnosis
- Has two GP practices as part of its portfolio of services, which include mental health practitioners and GPs, who are able to refer into the programme. The Carers Liaison Officer is due to undertake Carers Awareness training with GP staff, and GP Carers’ Champions will be launched in June 2018
- Will work to actively promote the Living Well Programme in both primary and secondary care settings

**DTHFT:**

Ensure that information around the living well programme is given to patients and carers on diagnosis and that information is also in the resource folder and on notice boards.

<table>
<thead>
<tr>
<th>10) Ensure the care and support needs of PLWD are met and regularly reviewed, as suggested in the Dementia Well Pathway</th>
<th><strong>STP Dementia &amp; Delirium Workstream:</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>The overarching key aims of the Derbyshire STP workstream for dementia and delirium are defined under the Dementia Well Pathway and there is a stated commitment by partners to ensure the needs of people with dementia are met and reviewed. The Strategy and Implementation partnerships will monitor this.</td>
<td></td>
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<tr>
<td><strong>DCC:</strong></td>
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<tr>
<td>DCC has adopted the Dementia Well Pathway as its framework for commissioning services for PLWD and their carers. This includes a commitment that all commissioned services will review needs regularly throughout the different stages of the pathway.</td>
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<tr>
<td><strong>CRH:</strong></td>
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<tr>
<td>The Trust undertakes a continuous patient experience survey to understand the experience of PLWD who use Trust services. In addition, friends and family feedback, concerns and complaints are regularly reviewed to identify any areas of good practice and opportunities for improvement.</td>
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<tr>
<td><strong>DTHFT:</strong></td>
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<tr>
<td>Support will be provided to the dementia key workers and dementia champions in the trust to ensure they can support all staff to ensure that the needs of PLWD are met and reviewed as required.</td>
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| 11) Ensure workers and professionals are able to identify patients/clients with dementia and their carers/family members, and have received adequate training to be able to meet their care and support needs in all settings | **STP Dementia & Delirium Workstream:**  
<table>
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<tbody>
<tr>
<td>The Delirium partnership within the STP workstream is currently working to produce a low or no cost training package around dementia and delirium for roll out across all partnership members to try and enable better awareness and response for PLWD, to prevent and manage this dangerous condition that PLWD are highly vulnerable to.</td>
</tr>
<tr>
<td><strong>Derbyshire CCGs:</strong></td>
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<tr>
<td>Encourage uptake of the DCC awareness sessions across Derbyshire.</td>
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<tr>
<td><strong>DCC:</strong></td>
</tr>
<tr>
<td>DCC welcome the positive feedback within the report regarding the Council’s own homecare staff support. DCC offers a specialist dementia training programme for care staff in addition to Care Certificate training, to ensure they work to the highest standards.</td>
</tr>
<tr>
<td><strong>CRH:</strong></td>
</tr>
<tr>
<td>The Carers Liaison Officer provides Carer Awareness training to staff. Champions will also be available to identify and support patients with dementia and their carers. Each carer seen by the</td>
</tr>
</tbody>
</table>
| 12) Maximise the need and importance of continuity, routine and training for homecare staff | **DCC:**  
- DCC is developing and expanding the Dementia Reablement Service, a short term specialist homecare service with a focus on continuity, routine and flexibility to help stabilise support at home  
- DCC aim to integrate the Dementia Reablement Service closely with the new NHS Dementia Rapid Response Teams, and share resources and training where possible  
- In May 2018 DCC will be holding a Best Practice in Dementia Care Conference, targeted at health and social care professionals, to enable further reflection and skill sharing when working with PLWD  
- The Carer’s Information, Advice and Support service provided by DCA now offers training and learning opportunities for carers, based on what carers have told us they would benefit from. |

| 13) Improve the experience of people involved with the DDSS in terms of more appropriate activities, more training for staff and ensure the one to one | **Derbyshire CCGs and DCC:**  
Derbyshire County Council and the NHS Derbyshire CCGs are in the process of re-procuring the DDSS. This will provide an opportunity to review and clarify the service priorities which include a high level of training for staff, a range of appropriate activity/peer support groups and accessible one to one support. |
Making Space/DDSS:

- **Appropriate activities**: Participants within all groups are encouraged to provide feedback following each session. This feedback is used to determine how future groups are planned, delivered and developed. Given the diversity of each group it can prove challenging to deliver sessions that are agreeable to everyone. DDSS make every effort to ensure that activity schedules are varied and appealing to participants.

- **Training for staff**: Many of the DDSS Dementia Advisors (DA’s) possess a wealth of knowledge amassed throughout long careers within a health and social care setting. For those less experienced there is regular, structured training aimed at improving and enhancing their knowledge base. All DAs have received dementia training from the Dementia Development Coordinator and Admiral Nurse Consultant for Making Space. There is a continual training structure in place covering a range of topics including: Person Centered Care, Distress Behaviour and Activity and Engagement. Although trained and knowledgeable, given the broad complexities surrounding dementia, there will be occasions where a DA will be required to carry out research in response to certain questions or requests for information. DAs will continue to receive regular training in order to maintain and develop knowledge and expertise. A package of training is being developed for the DDSS DAs by Stirling University, home of the Dementia Services Development Centre. This will be made available within the coming months.

- **One-to-one support**: DDSS have conducted almost 600 one-to-one support meetings with PLWD and their carers since October 2017. DDSS provides a service for anyone that wishes to receive one-to-one support from a DA. This applies to anyone that is new to the service and all existing service users, including those attending groups. DDSS will ensure that all group participants are aware of the option to receive one-to-one support outside of group settings. As part of a package of support developed during one-to-one meetings, referrals are made to other partner organisations including Derbyshire Carers Association to ensure that identified needs are met.

14) **Develop a role of a ‘Named Person’ to enable people to have one point of contact to be able to**

Derbyshire CCGs:

There are different models across the county, for example in Erewash the Community Connectors act as referrers and introducers to services as do the Local Area Coordinators in the City and the North.
| Access the needed advice and support | **DCC:**  
DCC will continue working with the service provider (Making Space) to ensure that all clients accessing the service have the contact details for a named worker.  

**CRH:**  
In our Primary Care Settings, the named person to enable people to have one point of contact to be able to access the advice and support needed would be the Care Coordinator. Within the Trust, advice and support is provided by the Enhanced Support, Older Person’s and DHCFT Liaison Teams; the Carers Liaison Officer acts as the named person for carers.  

**DTHFT:**  
The trust has a Dementia Lead Nurse who patients and relatives can be signposted to for advice and support.  

**Making Space/DDSS:**  
Upon referral to the DDSS a DA is allocated. This is determined by geographical location. This DA remains the single point of contact for a PLWD and their carer throughout their journey of support, and remains as such until all identified needs have been addressed. DDSS ensure that this is clearly communicated to anyone who is referred to the service, and that the relevant contact details are provided.  

| 15) Work to address the need for more support and activities for PLWD under the age of 65 | **STP Dementia & Delirium Workstream:**  
The STP will review key deliverables to make more explicit the commitment to people of all ages with dementia, and work within the implementation partnership to support partners in services designed for people below 65 years.  

**Derbyshire CCGs:**  
Through the Living Well Sessions. As part of the Derbyshire STP Dementia workstream DCC and other key partners are reviewing service access and support available for PLWD under the age of 65.  

**DCC:** |
Service user review panel groups, set up as part of the DFC project consider the need for a more diverse support and activity offer, linking with local DAAs in some areas of the county to facilitate new activities such as dementia-friendly swimming.

**CRH:**

Funding has recently been agreed for an activity room and Activity Coordinator, which will be available to all patients of all ages.

**DTHFT:**

The Trust has a link to the national dementia support manager for early onset dementia, who is able to offer support and advice to patients, carers and staff around early onset dementia. I will be meeting with the lead to discuss what activities are available for these patients to support them whilst an inpatient. We currently have numerous memory boxes that staff can utilise to provide distraction as well as use as activities for our patients. We also have resources we can use as activities which include using apps to listen to music, watch movies amongst various other activities.

**DCHS:**

DCHS is developing a Dementia Strategy which will include the following key strategic objective:

- Promotion of activities that improve the wellbeing of PLWD and their carers i.e. activities that promote physical and mental wellbeing, applicable to the services that we provide to PLWD.

**STP Dementia & Delirium Workstream:**

The STP fully recognises the importance of the carer, the carer’s needs and the carer’s own health as being key to the person with dementia and their health and well-being, and the STP partnership will work to support the integrated ways of working, including working with carer’s advocates and groups to ensure appropriate responses to the needs of carers. This action also relates to recommendations 17, 18 and 19.

**Derbyshire CCGs:**

We have been working with Derbyshire Carers regarding the feedback we had around carer’s assessment and support provision. The responsibility for this contract does sit with...
Derby City Council but Derbyshire Carers have taken up our offer for Person Centered Care training which helps them to have conversations focused on the carer’s needs and assets and their own wellbeing. Across the county there are also many discretionary voluntary and community groups and services funded to support carers. This includes day-to-day support for carers to go out, and also a carer’s retreat facility in Derby City - Haven House.

**DCC:**

Ensuring carers in Derbyshire have access to the support, advice and information to best meet their needs is one of the priorities outlined in the Council Plan. Therefore we will continue to ensure that services respond proactively to supporting carers. DCC has involved carers in co-producing services including the DDSS and Dementia Reablement Service and will continue to involve carers where possible.

**CRH:**

The Carers Liaison Officer is keen to ensure that the health and wellbeing of carers is supported within the role. The Carers Liaison Officer is able to offer one-to-one support and advice, as well as signposting to additional and future support services as appropriate, as are Carers Champions. The Carers Liaison Officer will continue to work with carers for up to two weeks after patient discharge; they will be referred to Derbyshire Carers after this time, should they require further support.

**DTHFT:**

All carers have access to John’s campaign room if they chose to stay with their loved ones, so they are able to take a break if needed. The facilities on offer are for carers to be able to access to sleeping facilities as well as kitchen facilities.

**Derbyshire Carers Association (DCA):**

- Some of this is the core approach with groups to get carers to focus on their needs as they often are so focused on the cared for. Keep healthy related topics are being introduced more into some of the support sessions, also looking at other areas in the country to see whether different models could be developed which offer carers specific keep healthy sessions, this will require joint working with health providers which may need commissioner support.
- Also commencing conversations with public health to see how the proposed PH hub model could also promote carer’s health
- Carer’s emergency card is an area which definitely needs more work and clarification about what it can and cannot do and what the expectations are from a carer’s view as well as the system. There is a lot of learning from the work so far especially in relation to how you can keep the info up to date, there is also the potential to link to the Summary Care record if commissioners can facilitate this so that Emergency Services and acute have a consistent approach.

### 17) Improve co-ordination of support services for carers, so they know where to go for advice, signposting and referrals

<table>
<thead>
<tr>
<th><strong>Derbyshire CCGs and DCC:</strong></th>
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<tbody>
<tr>
<td>As part of the Derbyshire STP Dementia workstream, DCC and other key partners will co-ordinate and integrate support services so they are more accessible and help to build carer’s confidence and resilience... A range of providers are working together within the Derbyshire STP Dementia workstream to identify opportunities for further co-ordination and integration of services. The views of carers of are particularly important as many PLWD are not able to express their views.</td>
</tr>
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**CRH:**

The Trust Carers Liaison Officer has established links with carers support services locally. The Trust’s Carers Working Group includes representation from Derbyshire Carers and other local agencies, as well as carer representatives.

**DTHFT:**

We have a dementia resource folder for families who require information and advice, which signposts them to various community groups where they are able to get advice and support.

**DCA:**

- We have become aware of these comments from carers and have in response looked at developing better joint working agreements/arrangements with our vol sector partners, especially Making Space. We feel this has the potential to be developed further, care needs to be taken not to duplicate or loose specialist knowledge
- DCA have now increased links with the Living Well Programme and with care co-ordinators
- There has been the promotion of the Carers in Derbyshire website, and DCA are improving their website over the next 6 months
- Over the next year there will be the development of the carer connect hubs
- In relation to carer’s assessment’s - Staff have had more training both in assessment and person centred approaches - this has taken place over the last 4 months.

**DCHS:**

DCHS is developing a Dementia Strategy which will include the following key strategic objectives:

- Development of a Carer Strategy which will support the carers and friends of PLWD. This will include carer assessments, advice and advocacy support
- Development of partnerships to improve collaborative working and improve integration of care. This will address the concerns raised in the report regarding the disjointed nature of services.

| 18) Ensure families who are self-funded have correct level of information and support | **DCC:**
| --- | --- |
|  | DCC continues to develop the Adult Care information offer for those who self-fund their care and support in line with the Care Act (2014). The Carers in Derbyshire website offers a range of vital information for self-funding families and we are committed to ensuring this information is accurate, up to date and relevant for self-funders.

**CRH:**

The Trust’s discharge team liaise with Social Services who will then visit patients and discuss their requirements. The Carers Liaison Officer is also able to support carers in finding information and support *(this also relates to recommendation 20).*

**DTHFT:**

We can signpost people to the relevant service *(Making Spaces)* who will be able to provide information and support to families.

| 19) Address the lengthy process for accessing respite care and to make it easier to access | **DCC:**
| --- | --- |
|  | DCC will review the information it provides regarding how to access respite care, to ensure this is accurate, up-to-date and easily accessible.

**DTHFT:**

The Lead Dementia Nurse has arranged a meeting with the
<table>
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<tr>
<th><strong>20) Provide clear information around Continuing Healthcare (CHC) and the use of jargon, so people are able to easily understand the process.</strong></th>
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<tbody>
<tr>
<td><strong>Discharge Lead in the Trust</strong> to discuss the process around people accessing respite care.</td>
</tr>
<tr>
<td><strong>Derbyshire CCGs:</strong></td>
</tr>
<tr>
<td>The CCGs have revised their website information on this. We are working with DCC, STP and other providers to ensure there is a targeted approach to ensure carers supporting those in receipt of fully funded CHC are proactively identified and supported to access the carer's service.</td>
</tr>
<tr>
<td><strong>DCC:</strong></td>
</tr>
<tr>
<td>DCC will continue to signpost people to free and independent advice such as that provided by Beacon (<a href="http://www.beaconhc.co.uk/">http://www.beaconhc.co.uk/</a> 0345 548 0300) regarding the Continuing Healthcare process.</td>
</tr>
<tr>
<td><strong>CRH:</strong></td>
</tr>
<tr>
<td>The Trust is keen that patients and carers understand the discharge process and utilises the ‘Red to Green’ process, which keeps patients and carers informed of what needs to happen prior to discharge.</td>
</tr>
<tr>
<td><strong>DTHFT:</strong></td>
</tr>
<tr>
<td>The Lead Dementia Nurse has arranged a meeting with the Discharge Lead in the Trust to discuss information around continuing health care to ensure that anyone needing to access it is understandable, and that jargon is at a minimum.</td>
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### Appendix 1

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<tr>
<th>Section</th>
<th>Description</th>
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| Preventing Falls | Measures to reduce falling risk include:  
- **Raised Bedroom Alarm**: Reduces the risk of falls at night by alerting caregivers when a fall occurs.  
- **Safety Bed Rails**: Provides support and limits the risk of falling from the bed.  
- **Non-slip Bed Sheets**: Reduces the risk of slipping on the bed.  
- **Soft Bed Pads**: Provides comfort and reduces the risk of pressure points.  
- **Bed Immobilization Devices**: Ensures the bed remains firm and reduces the risk of falling. |
| Supporting Wellbeing | Measures to support wellbeing include:  
- **Social Activities**: Engaging in social activities can help maintain mental and emotional health.  
- **Physical Exercise**: Regular exercise can improve physical health and reduce the risk of falls.  
- **Nutrition**: Eating a balanced diet can help maintain energy levels and reduce the risk of weakness.  
- **Hearing Aids**: Improves communication and reduces the risk of falls.  
- **Mobility Aids**: Provides additional support and reduces the risk of falls. |
| Supporting Carers | Measures to support carers include:  
- **Carer Support Services**: Provides support and reduces the risk of burnout.  
- **Carer Training**: Education and training can help carers to provide effective care.  
- **Carer Advice and Information**: Access to relevant information can help carers to make informed decisions.  
- **Carer Respite Care**: Provides a break for carers and reduces the risk of burnout.  
- **Carer Support Groups**: Provides a support network and reduces the risk of isolation. |
| Supporting Carers | Measures to support carers include:  
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- **Carer Support Groups**: Provides a support network and reduces the risk of isolation. |
11. Your feedback

Healthwatch Derbyshire is keen to find out how useful this report has been to you, and/or your organisation, in further developing your service. Please provide feedback as below, or via email.

1) I/we found this report to be: Useful / Not Useful

2) Why do you think this?

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3) Since reading this report:

a) We have already made the following changes: ........................................................
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b) We will be making the following changes: ........................................................
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Your name: .........................................................................................................................

Organisation: ......................................................................................................................

Email: .................................................................................................................................

Tel No: .................................................................................................................................

Please email to: karen@healthwatchderbyshire.co.uk or post to FREEPOST RTEE-RGYU-EUCK, Healthwatch Derbyshire, Suite 14 Riverside Business Centre, Foundry Lane, Milford, Belper, Derbyshire DE56 0RN.