Accelerated long-term forgetting in presymptomatic autosomal dominant Alzheimer’s disease: a cross-sectional study

Philip S J Weston, Jennifer M Nicholas, Susie M D Henley, Yuying Liang, Kirsty Macpherson, Elizabeth Donnachie, Jonathan M Schott, Martin N Rossor, Sebastian J Crutch, Christopher R Butler, Adam Z Zeman, Nick C Fox

Summary

Background Tests sensitive to presymptomatic changes in Alzheimer’s disease could be valuable for clinical trials. Accelerated long-term forgetting—during which memory impairment becomes apparent over longer periods than usually assessed, despite normal performance on standard cognitive testing—has been identified in other temporal lobe disorders. We assessed whether accelerated long-term forgetting is a feature of presymptomatic autosomal dominant (familial) Alzheimer’s disease, and whether there is an association between accelerated long-term forgetting and early subjective memory changes.

Methods This was a cross-sectional study at the Dementia Research Centre, University College London (London, UK). Participants were recruited from a cohort of autosomal dominant Alzheimer’s disease families already involved in research at University College London, and had to have a parent known to be affected by an autosomal dominant Alzheimer’s disease mutation, and not report any current symptoms of cognitive decline. Accelerated long-term forgetting of three tasks (list, story, and figure recall) was assessed by comparing 7-day recall with initial learning and 30-min recall. 7-day recognition was also assessed. Subjective memory was assessed using the Everyday Memory Questionnaire. The primary outcome measure for each task was the proportion of material retained at 30 min that was recalled 7 days later (ie, 7-day recall divided by 30-min recall). We used linear regression to compare accelerated long-term forgetting scores between mutation carriers and non-carriers (adjusting for age, IQ, and test set) and, for mutation carriers, to assess whether there was an association between accelerated long-term forgetting and estimated years to symptom onset (EYO). Spearman’s correlation was used to examine the association between accelerated long-term forgetting and subjective memory scores.

Findings Between Feb 17, 2015 and March 30, 2016, we recruited 35 people. 21 participants were mutation carriers (mean EYO 7·2 years, SD 4·5). Across the three tasks, we detected no differences between carriers and non-carriers for initial learning or 30-min recall. The proportion of material recalled at 7 days was lower in carriers than non-carriers for list (estimated difference in mean for list recall −30·94 percentage points, 95% CI −45·16 to −16·73; p=0·0002), story (−20·10, −33·28 to −6·91; p=0·0048), and figure (−15·41, −26·88 to −3·93; p=0·012) recall. Accelerated long-term forgetting was greater in carriers nearer to their estimated age at onset (p≤0·01 for all three tests). Mutation carriers’ 7-day recognition memory was also lower across all tasks (list [mean difference −5·80, 95% CI −9·96 to −2·47; p<0·01], story [−6·84, −10·94 to −3·37; p<0·01], and figure [−17·61, −27·68 to −7·72; p<0·01] recognition). Subjective memory scores were poorer in asymptomatic carriers compared with non-carriers (adjusted difference in means 7·88, 95% CI 1·36 to 14·41; p=0·016), and we found a correlation between accelerated long-term forgetting and subjective memory in mutation carriers.

Interpretation Accelerated long-term forgetting is an early presymptomatic feature of autosomal dominant Alzheimer’s disease, which appears to pre-date other amnestic deficits and might underpin subjective memory complaints in Alzheimer’s disease. Accelerated long-term forgetting testing might be useful in presymptomatic Alzheimer’s disease trials.

Funding MRC, NIHR, Alzheimer’s Research UK, Dementias Platform UK, Dunhill Medical Trust, ERUK, Great Western Research, Health Foundation, Patrick Berthoud Trust.

Copyright © The Author(s). Published by Elsevier Ltd. This is an Open Access article under the CC BY 4.0 license.

Introduction Pathological changes of Alzheimer’s disease develop years before clinical symptoms.1-3 The medial temporal lobe, which plays an important part in memory, is an early site of neurofibrillary tangle deposition and atrophy.1,4 Subtle cognitive impairment is likely to develop earlier than overt clinical symptoms, therefore cognitive measures sensitive to early changes would be valuable diagnostically and in presymptomatic trials. Autosomal dominant (familial) Alzheimer’s disease, due to mutations in presenilin 1 (PSEN1), presenilin 2 (PSEN2), or amyloid precursor protein (APP) genes,
Research in context

Evidence before this study
Accelerated long-term forgetting is a feature of temporal lobe epilepsy. We investigated whether accelerated long-term forgetting had been assessed in early Alzheimer’s disease by searching PubMed for articles published between Jan 1, 1980, and April 11, 2016, with the terms “Alzheimer’s disease” AND “long-term forgetting” OR “accelerated forgetting”. No articles that reported accelerated long-term forgetting in presymptomatic Alzheimer’s disease were found. We then did a second PubMed search using the terms “Alzheimer’s disease” AND “familial” OR “autosomal dominant” AND “presymptomatic” OR “preclinical” AND “cognitive” OR “memory” to identify studies of early cognitive change in autosomal dominant Alzheimer’s disease. No language restrictions were used in either of the searches. The general pattern reported in these studies was of episodic memory decline evident up to 2–3 years before the expected onset of symptomatic disease, although one longitudinal study showed measurable decline only after the time of estimated onset. No studies assessing episodic memory had a retention interval of longer than 30 min. Visual short-term memory binding and dual task performance also declined in mutation carriers around 2 years before symptom onset. No studies were identified that showed earlier presymptomatic cognitive decline.

Added value of this study
To our knowledge, this study presents the first assessment of long-term forgetting in asymptomatic individuals who perform normally on conventional memory tests but have known Alzheimer’s pathology. We found that differences in accelerated long-term forgetting were detectable in individuals who were on average 7 years away from predicted onset of symptomatic disease. We found that long-term retention of both verbal and visual material was affected, with impairment of recall and recognition. Accelerated long-term forgetting appears to pre-date otherwise previously identified early Alzheimer’s disease-related cognitive changes. Subtle increases in subjective cognitive concerns are also present several years before expected symptom onset.

Implications of all the available evidence
Our results add to existing evidence that episodic memory is impaired before symptom onset in individuals with Alzheimer’s disease pathology. However, by assessing memory retention over a longer interval than is conventionally used, it is possible to detect verbal and visual memory deficits earlier than previously thought. Accelerated long-term forgetting appears to be one of the earliest detectable features of Alzheimer’s disease-related cognitive decline. The results of this study extend our understanding of the temporal sequence of memory decline in Alzheimer’s disease; it was previously possible to separate healthy ageing from Alzheimer’s disease dementia and mild cognitive impairment by testing encoding and short interval retention. Assessment of accelerated long-term forgetting appears to provide a means of discriminating healthy ageing from asymptomatic individuals in the presymptomatic phase of Alzheimer’s disease. Our findings of early, subtle subjective memory changes before overt symptoms in individuals with known Alzheimer’s disease pathology, support research suggesting that subjective changes are predictive of future decline. The potential association between subjective memory change and accelerated long-term forgetting is consistent with previous epilepsy studies and, if confirmed, might explain the mechanism of early subjective change in Alzheimer’s disease. Future work is required to define more precisely when deficits appear relative to actual symptom onset and to determine the use of accelerated long-term forgetting assessment in presymptomatic Alzheimer’s disease trials.

shares many features, both pathophysiological and clinically, with the much more common sporadic form of the disease. Autosomal dominant Alzheimer’s disease mutation carriers have relatively predictable ages at symptom onset based on family history, and therefore provide an opportunity to study presymptomatic cognitive change where time to symptom onset can be estimated.

Deficits in episodic memory are one of the earliest detectable cognitive abnormalities in Alzheimer’s disease. However, there is a substantial delay between pathological change and measurable cognitive decline. Accelerated long-term forgetting is a form of memory impairment that has been described in patients with temporal lobe epilepsy. It refers to a process whereby new material appears to be encoded and retained normally over periods of up to 30 min—consistent with normal performance on standard memory tests—but is then forgotten at an abnormally rapid rate over the following hours to weeks. Accelerated long-term forgetting might represent disruption in memory consolidation, which refers to gradual post-acquisition stabilisation of long-term memories, with the medial temporal lobe known to play an important part.

Accelerated long-term forgetting was associated with subjective cognitive concerns in patients with epilepsy. Such concerns might also be a preclinical harbinger of cognitive decline in Alzheimer’s disease.

Accelerated long-term forgetting has recently been assessed in a mouse model of presymptomatic autosomal dominant Alzheimer’s disease. Genetically modified mice had abnormal memory retention after 7 days, despite normal learning and retention over shorter intervals.

We hypothesised that accelerated long-term forgetting would also be a feature of presymptomatic autosomal dominant Alzheimer’s disease in human beings, with mutation carriers forgetting more material over 7 days.
compared with non-carriers, despite similar learning and short interval (30 min) retention. Additionally, we investigated whether accelerated long-term forgetting is associated with subjective cognitive concerns.

Methods
Study design and participants
We did a cross-sectional study at the Dementia Research Centre, University College London (London, UK). Participants were recruited from families who were already actively involved (ie, consented to inclusion) in a long-running longitudinal study of autosomal dominant Alzheimer’s disease at University College London. All participants, by having an affected parent, were at 50% risk of carrying a pathogenic autosomal dominant Alzheimer’s disease mutation.² Participants were also required to be asymptomatic, with neither they nor their informant reporting progressive cognitive symptoms. A further inclusion criterion was being aged older than 18 years. Participants were excluded from the study if they had substantial co-existing neurological or psychiatric disease. The study was approved by the local Research Ethics Committee and all participants provided written informed consent.

Procedures
Genetic testing was done to establish mutation status and results were provided only to statisticians, ensuring that participants and clinicians assessing them remained blind to genetic status. All individuals identified a close informant who was interviewed separately to gain a collateral history. Estimated years to symptom onset were calculated by subtracting the participant’s age at testing from the age at which their affected parent first developed progressive cognitive symptoms.

Participants underwent neurological examination and assessment using the mini mental state examination and the Clinical Dementia Rating scale,²⁶ which incorporates information from both participant and informant on day-to-day cognition. Neuropsychological testing included measures of general intellectual functioning (Wechsler Abbreviated Scale of Intelligence [WASI]),²¹ verbal and visual recognition memory (Recognition Memory Test for words and faces),²² and paired associate learning (Camden paired associate learning test).²³ Depression and anxiety were assessed using the Hospital Anxiety and Depression Scale (HADS).²⁸ Subjective memory was assessed using the Everyday Memory Questionnaire (EMQ),²⁹ scored between 0 and 90, with higher scores indicating greater concerns.

Long-term forgetting was assessed using test materials from the Adult Memory and Information Processing Battery.²⁶ Assessment was 1–3 months after background clinical data collection. This 2-month window was allowed because of practical considerations that meant it was not always possible for every participant to return for assessment after exactly the same interval. A minimum 1-month gap was left between neuropsychological testing and long-term forgetting assessment so that participants did not confuse material from the two types of test. Participants were assessed on three tests: learning and recall of a 15-item word list, a short story, and a complex visual figure. For the word list, participants had to learn the material to a minimum required accuracy of 80% over a minimum of four and maximum of ten trials. For the story, the minimum required accuracy was 80%, over a minimum of two and maximum of ten trials. For the figure, participants were first asked to copy the figure as accurately as possible on to a separate piece of paper; the figure and copy were then removed and participants were asked to draw the figure again from memory. Participants were then tested on free recall of the word list, story, and figure 30 min after presentation of the last learning trial for each test. All three tests (list, story, and figure) were marked by the same assessor (PSJW) using validated marking criteria.²⁶

There were two versions of each test. Participants were randomly assigned to either set one or set two (a random number was generated [either 1 or 2] before testing each participant, which was then used to decide the set), except for families in which two members were participating, in which case the second participant in a family was always assessed on a different version from the first participant. Participants were requested not to discuss the tests with other participants.

Following assessment, participants were given an envelope (appendix), which they were asked not to open until told to do so during a telephone call 7 days later. Participants were not told that the memory tests would be repeated. At the 7-day telephone call, participants’ free recall of the test materials was reassessed. For the figure, they were asked to draw the figure on a blank piece of paper included in the envelope, and for the verbal tests recall was done orally. Forced-choice recognition memory was then assessed. For the word list, participants were read 15 semantically unrelated word pairs, and asked to identify which of each pair was in the original list. A similar three-alternative forced-choice response procedure was done for 12 separate aspects of the story. For the figure, participants were asked to view a sheet (contained in a separate envelope) showing four sets of three similar

| Table 1: Participant characteristics |
| ----------------------------------- |
| **Mutation carriers (n=21)** | Non-carriers (n=14) |
| **Sex** | | |
| Male | 11 (52%) | 6 (43%) |
| Female | 10 (48%) | 8 (57%) |
| **Age (years)** | 38 0 (5.9) | 39 2 (7.9) |
| **Estimated years to symptom onset** | 7 2 (4.5) | NA |
| **Years of education** | 13 8 (2.5) | 14 4 (2.2) |

Data are n (%) or mean (SD). NA—not applicable.
illustrations and were asked to mark which of each set of three exactly matched part of the figure. Figure assessments were returned in a stamped addressed envelope.

Outcomes

The primary outcome measure of interest was the proportion of material (list, story, and figure) retained at 30 min that was recalled 7 days later (ie, 7-day recall divided by 30-min recall). This approach, used in previous studies in patients with epilepsy, allows comparison of group differences at 7 days, adjusting within each individual for short interval (30-min) forgetting. Secondary outcomes were 7-day recall and recognition scores for each of the three tasks. The only other secondary outcome was subjective memory score, although this was separate from the long-term forgetting analysis.

Statistical analysis

Scores on standard cognitive tests, HADS, EMQ, and accelerated long-term forgetting initial learning (score on the final learning trial), number of trials to criterion, 30-min recall, 7-day recall, 7-day score divided by 30-min score, and 7-day recognition were compared between mutation carriers and non-carriers using linear regression with robust SEs. All tests were performed with robust SEs accounting for clustering of participants within families. Secondary outcomes for each of the three tests were 7-day recall and recognition scores, with the primary outcome measure being 7-day recall divided by 30-min recall. For the list and story, learning score refers to the score from the final learning trial.

Statistical analysis

Data are median (IQR); p values are shown for variables that satisfied the assumptions of the linear regression model. For other variables, parametric assumptions were not met and therefore p<0.05 or p>0.05 was inferred from bootstrapped 95% CIs. Differences in means, proportions, or medians were compared using twotailed Student’s t tests for continuous data, the two-tailed Fisher exact test for categorical data, and the one-tailed Wilcoxon rank sum test for ordinal data. For data with non-normal distributions, statistical inference was based on non-parametric tests. If plots of model residuals indicated a departure from the assumption of normal distribution with constant variance, robust SEs were used. For variables without such a distribution, for which robust SEs were not appropriate, statistical inference was based on non-parametric tests.
bias-corrected and accelerated 95% CIs and 99% CIs from 10 000 bootstrap replications clustered on family.

Comparison of HADS, mini mental state examination, Recognition Memory Test faces, Recognition Memory Test words, Camden paired associated learning test, and EMQ was adjusted for age and intelligence quotient (IQ; WASI total). Comparison of WASI performance IQ and WASI verbal IQ was unadjusted, as these measures were corrected for age. Comparison of groups with regards to accelerated long-term forgetting scores was adjusted for age, IQ, and test set.

Non-parametric receiver operating characteristic (ROC) curves were used to quantify the ability of the 7-day score divided by the 30-min score on each of the accelerated long-term forgetting tests to discriminate between carriers and non-carriers. Areas under the curves (AUCs) were calculated and cutoffs, specificities, and sensitivities reported.

The association between EMQ and 7-day score divided by 30-min score on each accelerated long-term forgetting test was assessed using the non-parametric Spearman correlation coefficient, with inference based on non-parametric bias-corrected and accelerated 95% and 99% CIs from 10 000 bootstrap replications clustered on family. For analyses involving EMQ, only participants who were unaware of their mutation status were included, as knowledge of status might bias an individual’s perception of their memory.

Results were interpreted taking into consideration that a Bonferroni correction for use of three accelerated long-term forgetting tests (list, story, figure) would require p<0·017 (ie, 0·05 divided by 3) for formal statistical significance.

For the four figure recall and recognition assessment documents that were not returned, these data were assumed to be missing entirely at random. Data were not imputed for missing cases.

All statistical analyses were done using Stata version 14.0 or later.

Role of the funding source
The funders of the study had no role in study design, data collection, data analysis, data interpretation, or writing of the report. The corresponding author had full access to all the data in the study and had final responsibility for the decision to submit for publication.

Results
We recruited 35 asymptomatic individuals from 19 autosomal dominant Alzheimer’s disease families between Feb 17, 2015, and March 30, 2016. Nine participants (five mutation-positive, four mutation-negative) had previously chosen to have clinical predictive genetic testing, and so were aware of their genetic status. 21 (60%) of 35 participants were mutation carriers. All participants learned the word list and story to the required 80% accuracy within the allowed number of trials. Across all three accelerated long-term forgetting tests, adjusting for age, IQ, and test set, and accounting

![Figure 1: Long-term forgetting assessments](image-url)

Scores for (A) word list, (B) story, and (C) figure were adjusted for age, intelligence quotient (WASI total score), and test set. Robust SEs accounted for within-family clustering. WASI=Wechsler Abbreviated Scale of Intelligence.
for clustering within families, the ability of mutation carriers to learn the material did not differ from non-carriers, both in terms of initial score and in the number of learning trials (table 3). Recall score at 30 min for all three tasks did not differ between groups (table 3, figure 1). All participants completed all assessments. However, four participants failed to return the 7-day figure recall and recognition assessment documents, so these could not be included in the analysis of these measures.

For 7-day recall scores divided by 30-min recall scores, mutation carriers did worse than non-carriers for list (adjusted difference in means –30·94 percentage points, 95% CI –45·16 to –16·73; p=0·0002), story (–20·10, –33·28 to –6·91; p=0·0048) and figure (–15·41, –26·88 to –3·93; p=0·012) recall.

The poorer retention of material at 7 days in mutation carriers was also seen in lower mean 7-day recall scores for list and story categories than in non-carriers, but not for figure (table 3). Mutation carriers also did worse than non-carriers on 7-day list, story, and figure recognition (table 3).

ROC analysis for discrimination between mutation carriers and non-carriers for 7-day score divided by 30-min score revealed an AUC of 0·86 (95% CI 0·66–0·95) for list, 0·84 (0·62–0·96) for story, and 0·79 (0·53–0·93) for figure scores (figure 2). A cutoff of less than 60% on list long-term retention gave 76% sensitivity and 71% specificity for mutation carriers versus non-carriers. A cutoff of less than 88% for story recall gave 90% sensitivity and 71% specificity. A cutoff of less than 75% for figure recall gave 79% sensitivity and 67% specificity.

There was an association between long-term forgetting and EYO for list, story, and figure scores, with the severity of accelerated long-term forgetting increasing with proximity to predicted symptom onset (figure 3).

Neither mutation carriers (median 18·5 [IQR 12–23] of 90 maximum) nor non-carriers (12·5 [8–15] of maximum of 90) had high EMQ scores (having excluded participants who knew their genetic status). However, mutation carriers had more subjective memory concerns than did non-carriers (adjusted difference in means 7·88, 95% CI 1·36–14·41; p=0·016) (figure 4). In mutation carriers, an association existed between higher EMQ and poorer long-term retention (7-day recall score divided by 30-min recall score), which reached statistical significance for both list and story, but not for figure (figure 5). There was no evidence of an association between EMQ score and performance on standard cognitive tests.

Discussion
We found that accelerated long-term forgetting was a feature of presymptomatic autosomal dominant Alzheimer’s disease in a group of individuals who were on average 7 years from estimated symptom onset. Despite similar learning and ability to recall over a short interval, reassessment 7 days later showed that
presymptomatic mutation carriers had forgotten more than had non-carriers. This finding was consistent for recall and recognition memory, and for verbal and non-verbal material. Among mutation carriers, severity of accelerated long-term forgetting increased with proximity to estimated symptom onset.

By contrast, mutation carriers showed no impairment in performance on conventional memory tests, which included both recognition memory for words and faces and paired associate learning. This finding suggests that differences in long-term forgetting are unlikely to be due to problems with encoding or early retention, but rather are caused by impairment of long-term memory consolidation. Although impairment of episodic memory is recognised as a central component of the Alzheimer’s disease phenotype, relatively little is known about how long-term consolidation of memory is affected. Some studies have investigated potential consolidation deficits in early Alzheimer’s disease, but it is difficult to measure long-term retention of memory reliably and compare a disease group with controls, unless the groups are equal for initial encoding and short interval retention, which in these studies was not the case.

By recruiting a presymptomatic Alzheimer’s disease cohort, and testing retention over a longer period than in standard neuropsychological assessment, we showed that impairment of long-term retention is an early amnestic feature of Alzheimer’s disease, and might be one of the earliest features of Alzheimer’s disease-related cognitive decline.

A previous study of autosomal dominant Alzheimer’s disease tested recognition memory over a more conventional interval of about 5 min and identified presymptomatic memory changes, but only 1–2 years before symptom onset. Other studies of presymptomatic carriers (from the PSEN1 Colombian kindred) showed reduced cognitive performance in several domains, including visual short-term memory binding, dual task performance, and lexical-semantic processing; however, participants were closer to expected symptom onset than were those in our study. Another study of the same cohort found that earlier decline could be identified by using a composite score comprising multiple tests. A similar composite cognitive test approach, which included paired associate learning, delayed recall, naming, and digit-symbol substitution...
(an assessment of short-term memory and processing speed) identified decline in individuals in the late stage of presymptomatic sporadic Alzheimer’s disease.35 We also found group differences in recognition memory at 7 days. Impaired recognition, which implies a so-called storage deficit rather than an isolated retrieval deficit, is consistent with pathology primarily affecting the medial temporal lobe,36 a brain region affected early in Alzheimer’s disease.

ROC analysis indicated that accelerated long-term forgetting assessments might provide a means of discriminating between presymptomatic Alzheimer’s disease and healthy ageing, with discriminatory power being stronger for verbal tests (list and story) than for the figure test. However, the sample size of the current study limits the precision with which AUCs can be estimated. Moreover, specificity values for our proposed cutoffs are suboptimal, and further work is needed to determine the most suitable cutoffs.

Our findings advance existing understanding of the stages of memory decline in Alzheimer’s disease. Previous studies37 have reported discrimination of Alzheimer’s disease dementia from Alzheimer’s disease mild cognitive impairment based on initial encoding of memory, which has been found to be impaired in the dementia stage but not in mild cognitive impairment. Similarly, Alzheimer’s disease mild cognitive impairment can be separated from presymptomatic disease and healthy ageing based on short-term retention (eg, over 30 min), which is impaired only in Alzheimer’s disease mild cognitive impairment. Our results suggest that incorporating tests of long-term (eg, 7-day) retention might allow individuals with presymptomatic Alzheimer’s disease (but without conventional deficits) to be distinguished from healthy ageing (table 4).

Other than accelerated long-term forgetting, the only group difference where carriers performed worse than non-carriers on cognitive testing was for performance IQ. This finding is consistent with previous findings in autosomal dominant Alzheimer’s disease that, along with episodic memory, reduced performance IQ is one of the earliest cognitive changes.3 However, the proportional difference in performance IQ between groups was much smaller than for accelerated long-term forgetting. As we adjusted for WASI score in all regression analyses, it is unlikely that the difference in performance IQ influenced the main study findings.

Despite reporting no overall concerns about their memory and being unaware of their genetic status, we found that presymptomatic mutation carriers had worse subjective memory scores than non-carriers, which supports emerging evidence that subtle subjective cognitive concerns might be a harbinger of Alzheimer’s disease-related cognitive decline.19–20 In mutation carriers, greater subjective cognitive concerns were associated with accelerated long-term forgetting for two of the three components (list and story). This association is consistent with findings from previous epilepsy studies,11,12 although a study13 assessing a combination of different neurological diagnoses did not identify a relationship. Early subjective memory concerns in Alzheimer’s disease, in the absence of conventional deficits, might reflect accelerated forgetting and impairments in long-term recall.

Our findings have parallels with a study of APP transgenic mice.37 Very young mice that had normal learning of a spatial navigation task and normal retention over a short interval had significantly reduced retention compared with wild-type mice when retested 7 days later. Furthermore, the deficit in long-term retention could potentially be rescued by anti-amyloid β immunotherapy.38 These results, alongside our findings, suggest that assessment of long-term forgetting in human beings might be useful as a marker of treatment response in presymptomatic therapeutic trials.

Our study has several limitations. First, the sample size is small, primarily because of the relative rarity of autosomal dominant Alzheimer’s disease. Replication of our findings in both familial and sporadic presymptomatic Alzheimer’s disease cohorts is therefore important. Although participants were asked not to discuss the testing following the initial learning and 30-min recall task, some individuals could have rehearsed the material during the intervening period, which is an inherent difficulty with accelerated long-term forgetting testing.39 However, when asked following the 7-day testing, all participants said that they had not rehearsed. Rehearsal might be more difficult to avoid in a trial with repeated measures, which could reduce test sensitivity. Therefore, it is important to identify methods to reduce the potential for rehearsal, such as embedding the accelerated long-term forgetting assessment material among other unrelated cognitive tests or making use of recognition tests for material that is difficult to rehearse.40–42 We found a ceiling effect on the 7-day recognition testing, which could be reduced in future studies by using more demanding tests (eg, paired associate learning).42

We did not assess participants’ sleep between initial learning and 7-day testing. Sleep is thought to play an

| Memory encoding | Early retention | Long-term retention |
|-----------------|----------------|--------------------|
| Alzheimer’s disease dementia | Reduced | Reduced | Reduced |
| Mild cognitive impairment | Normal/unchanged | Reduced | Reduced |
| Presymptomatic Alzheimer’s disease | Normal/unchanged | Normal/unchanged | Reduced |
| Healthy ageing | Normal/unchanged | Normal/unchanged | Normal/unchanged |

Previous studies have established differences in memory function between Alzheimer’s disease dementia, Alzheimer’s disease mild cognitive impairment, and healthy ageing, based on tests of memory encoding and retention over a short interval. The results of our study allow us to propose a third column, for long-term retention, which shows impairment several years before onset of noticeable symptoms.
important part in memory consolidation." However, a previous epilepsy study found sleep quality to have no direct effect on accelerated long-term forgetting, and we found no group difference in mood, which correlates with sleep. A 7-day assessment was done remotely, which allows other sources of variability, including the surrounding environment, to add noise. Nevertheless, we detected group differences in retention at 7 days, suggesting robust effects.

We used parental age at onset to estimate when an individual would develop Alzheimer’s disease symptoms. Although parental age at onset correlates closely with actual age at onset, it is a proxy measure, and only with longer-term follow-up can actual ages at symptom onset be confirmed.

In summary, we showed that accelerated long-term forgetting is present in autosomal dominant Alzheimer’s disease mutation carriers, who were on average 7 years from predicted symptom onset, and thus might be a very early feature of Alzheimer’s disease-related cognitive decline. Subtle increased subjective cognitive concerns also appear to be a presymptomatic feature of autosomal dominant Alzheimer’s disease, which could be underpinned by accelerated long-term forgetting. Studies of accelerated long-term forgetting might provide insights into very early cognitive impairments in Alzheimer’s disease, and accelerated long-term forgetting might be useful as part of inclusion criteria or as an outcome measure in presymptomatic Alzheimer’s disease trials.

Contributors
CRB, AZZ, NCF, and SJC conceived the study. PSJW, YL, and SMDH designed the accelerated long-term forgetting testing framework. PSJW collected all clinical and accelerated long-term forgetting testing data. KM and ED collected standard neuropsychological test data. JMN led the statistical analysis. PSJW, JMW, SMDH, MNR, JMS, SJC, CRB, AZZ, and NCF interpreted the data. PSJW drafted the initial manuscript. All authors contributed to revision and editing of the manuscript.

Declaration of interests
JMS has received research funding from AVID Radiopharmaceuticals (a wholly owned subsidiary of Eli Lilly), has consulted for Roche Pharmaceuticals and Eli Lilly, and serves on a Data Safety Monitoring Committee for Axon Neuroscience. NCF reports fees (paid to University College London) for consultancy from Eli Lilly, Novartis, Sanofi, Roche/Genentech, and GlaxoSmithKline, and for serving on a Data Monitoring Committee for Biogen. All other authors declare no competing interests.

Acknowledgments
We thank the participants and their families for their generous support of this study. We are grateful to Michaela Dewar who developed some of the test materials used as filler tasks between the accelerated long-term forgetting learning and 30-min recall assessments. PSJW is supported by an MRC Clinical Research Training Fellowship. SJC is supported by an Alzheimer’s Research UK Senior Investigator Fellowship. CRB holds an MRC Clinician Scientist fellowship. The work of AZZ and CRB has been principally supported by the Dunhill Medical Trust, Epilepsy Research UK, Great Western Research, the Health Foundation, and the Patrick Berthoud Trust. The Dementia Research Centre is an Alzheimer’s Research UK Coordinating Centre and is grateful for support from the NIHR Queen Square Dementia Biomedical Research Unit, the MRC Dementias Platform UK, the Leonard Wolfson Experimental Neurology Centre, and the UK Dementia Research Institute. The study was done at the Dementia Research Centre, University College London Institute of Neurology (London, UK) who received a proportion of funding from the Department of Health’s NIHR Biomedical Research Centres funding scheme.

References
1. Bateman RJ, Xiong C, Benzinger TL, et al. Clinical and biomarker changes in dominantly inherited Alzheimer’s disease. N Engl J Med 2012; 367: 795–804.
2. Villemagne VL, Burnham S, Bourgeat P, et al. Amyloid β deposition, neurodegeneration, and cognitive decline in sporadic Alzheimer’s disease: a prospective cohort study. Lancet Neurol 2013; 12: 357–67.
3. Braak H, Braak E. Evolution of neuronal changes in the course of Alzheimer’s disease. J Neural Transm Suppl 1998; 53: 127–40.
4. Brier MR, Gordon B, Friedrichsen K, et al. Tau and Aβ imaging, CSF measures, and cognition in Alzheimer’s disease. Sci Transl Med 2016; 8: 338ra66.
5. Ridha BH, Barnes J, Bartlett JW, et al. Tracking atrophy progression in familial Alzheimer’s disease: a serial MR image study. Lancet Neurol 2006; 5: 828–34.
6. Bateman RJ, Aiyesen PS, De Strooper B, et al. Autosomal-dominant Alzheimer’s disease: a review and proposal for the prevention of Alzheimer’s disease. Alzheimers Res Ther 2011; 3: 1.
7. Rymdaz DC, Acosta-Baena N, Aiyesen PS, et al. Symptom onset in autosomal dominant Alzheimer disease: a systematic review and meta-analysis. Neurology 2014; 83: 253–60.
8. Fox NC, Warrington EK, Seffier AI, Agnew SK, Rossor MN. Presymptomatic cognitive deficits in individuals at risk of familial Alzheimer’s disease. A longitudinal prospective study. Brain 1998; 121: 1631–39.
9. Yau WW, Tudorascu DL, McDade EM, et al. Longitudinal assessment of neuroimaging and clinical markers in autosomal dominant Alzheimer’s disease: a prospective cohort study. Lancet Neurol 2015; 14: 804–13.
10. Linn RT, Wolf PA, Bachman DL, et al. The ‘preclinical phase’ of probable Alzheimer’s disease. A 13-year prospective study of the Framingham cohort. Arch Neurol 1995; 52: 485–90.
11. Blake RV, Wroe SJ, Breen EK, McCarthy RA. Accelerated forgetting in patients with epilepsy: evidence for an impairment in memory consolidation. Brain 2000; 123: 472–83.
12. Butler CR, Graham KS, Hodges JR, Kapur N, Wardlaw JM, Zeman AZ. The syndrome of transient epileptic amnesia. Ann Neurol 2007; 61: 367–98.
13. Hoeffeizers S, Dewar M, Della Sala S, Zeman A, Butler C. Accelerated long-term forgetting in transient epileptic amnesia: an acquisition or consolidation deficit? Neuropsychologia 2013; 51: 1549–55.
14. Duda Y, Karmi A, Born J. The consolidation and transformation of memory. Neuron 2015; 88: 20–32.
15. Dik MG, Jonker C, Comijs HC, et al. Memory complaints and APOE ε4 accelerate cognitive decline in cognitively normal elderly. Neurology 2001; 57: 2237–22.
16. Reisberg B, Prichep I, Mosconi L, et al. The pre-mild cognitive impairment, subjective cognitive impairment stage of Alzheimer’s disease. Alzheimer’s Dement 2008; 4 (suppl 1): 598–108.
17. Jessen F, Amarinego RF, van Bostel M, et al. A conceptual framework for research on subjective cognitive decline in preclinical Alzheimer’s disease. Alzheimers Dement 2014; 10: 844–52.
18. Perrotin A, La Joie R, De La Sayette V, et al. Subjective cognitive decline in cognitively normal elders from the community or from a memory clinic: differential affective and imaging correlates. Alzheimers Dement 2017; 13: 550–60.
19. Begloupolou V, Tullioch J, Roe AD, et al. Early detection of cryptic memory and glucose uptake deficits in pre-pathological APP mice. Nat Commun 2016; 7: 11761.
20. Morris JC. The Clinical Dementia Rating (CDR): current version and scoring rules. Neurology 1993; 43: 2412–14.
21. Wechsler Abbreviated Scale of Intelligence (WASI). San Antonio, TX: Psychological Corporation, 1999.
22. Recognition memory test. Windsor, UK: nferNelson, 1984.
23. The Camden Memory Tests. Howe: Psychology Press, 1996.
24. Zigmond AS, Snaith RP. The hospital anxiety and depression scale. Acta Psychiatr Scand 1983; 67: 361–70.
25 Thompson PJ, Corcoran R. Everyday memory failures in people with epilepsy. Epilepsia 1992; 33 (suppl 6): S18–20.

26 The adult memory and information processing battery (AMIPB): test manual. Leeds: A K Coughlin, Psychology Dept, St James’ Hosp, 1985.

27 Butler CR, Bhaduri A, Acosta-Cabronero J, et al. Transient epileptic amnesia: regional brain atrophy and its relationship to memory deficits. Brain 2009; 132: 357–68.

28 Koivunen J, Karrasch M, Scheinin NM, et al. Cognitive decline and amyloid accumulation in patients with mild cognitive impairment. Dement Geriatr Cogn Disord 2012; 34: 31–37.

29 Elliott G, Isaac CI, Muhlert N. Measuring forgetting: a critical review of accelerated long-term forgetting studies. Cortex 2014; 54: 16–32.

30 Tort-Merino A, Valech N, Péhaloza C, et al. Early detection of learning difficulties when confronted with novel information in preclinical Alzheimer’s disease stage 1. J Alzheimers Dis 2017; 58: 855–70.

31 MacPherson SE, Parra MA, Moreno S, Lopera F, Della Sala S. Dual task abilities as a possible preclinical marker of Alzheimer’s disease in carriers of the E280A presenilin-1 mutation. J Int Neuropsychol Soc 2012; 18: 214–41.

32 Parra MA, Abrahams S, Logie RH, Méndez LG, Lopera F, Della Sala S. Visual short-term memory binding deficits in familial Alzheimer’s disease. Brain 2000; 133: 2702–13.

33 Arango-Lasprilla JC, Cueto F, Valencia C, Uribe C, Lopera F. Cognitive changes in the preclinical phase of familial Alzheimer’s disease. J Clin Exp Neuropsychol 2007; 29: 892–900.

34 Ayutyanont N, Langbaum JB, Hendrix SB, et al. The Alzheimer’s prevention initiative composite cognitive test score: sample size estimates for the evaluation of preclinical Alzheimer’s disease treatments in presenilin 1 E280A mutation carriers. J Clin Psychiatry 2014; 75: 652–60.

35 Soldan A, Pettigrew C, Gai Q, et al. Hypothetical preclinical Alzheimer disease groups and longitudinal cognitive change. JAMA Neurad 2016; 73: 698–705.

36 Simons JS, Spiers HJ. Prefrontal and medial temporal lobe interactions in long-term memory. Nat Rev Neurosci 2003; 4: 637–48.

37 Ally BA, Hussey EP, Ko PC, Molitor RJ. Pattern separation and pattern completion in Alzheimer’s disease: evidence of rapid forgetting in amnestic mild cognitive impairment. Hippocampus 2013; 23: 1246–58.

38 Manes F, Serrano C, Calcagno ML, Cardozo J, Hodges J. Accelerated forgetting in subjects with memory complaints. A new form of mild cognitive impairment? J Neurol 2008; 255: 1067–70.

39 van der Werf SP, Geurts S, de Ward MM. Subjective memory ability and long-term forgetting in patients referred for neuropsychological assessment. Front Psychol 2016; 7: 605.

40 Bateman RJ, Benzinger TL, Berry S, et al. The DIAN-TU Next Generation Alzheimer’s disease prevention trial: adaptive design and disease progression model. Alzheimers Dement 2017; 13: 8–19.

41 Dewar M, Alber J, Cowan N, Della Sala S. Boosting long-term memory via wakeful rest: intentional rehearsal is not necessary, consolidation is sufficient. PLoS One 2014; 9: e109542.

42 Blackwell AD, Sahakian BJ, Vesey R, Semple JM, Robbins TW, Hodges JR. Detecting dementia: novel neuropsychological markers of preclinical Alzheimer’s disease. Dement Geriatr Cogn Disord 2004; 17: 42–48.

43 Rasch B, Born J. About sleep’s role in memory. Physiol Rev 2013; 93: 681–766.

44 Atherton KE, Nobre AC, Zeman AZ, Butler CR. Sleep-dependent memory consolidation and accelerated forgetting. Cortex 2014; 54: 92–105.

45 Boivin DB, Czeisler CA, Djik DD, et al. Complex interaction of the sleep-wake cycle and circadian phase modulates mood in healthy subjects. Arch Gen Psychiatry 1997; 54: 145–52.