BRAIN COMMUNICATIONS

Single-subject grey matter network trajectories over the disease course of autosomal dominant Alzheimer's disease

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The investigators of Dominantly Inherited Alzheimer Network (DIAN) are given in the Appendix I section.

Structural grey matter covariance networks provide an individual quantification of morphological patterns in the brain. The network integrity is disrupted in sporadic Alzheimer's disease, and network properties show associations with the level of amyloid pathology and cognitive decline. Therefore, these network properties might be disease progression markers. However, it remains unclear when and how grey matter network integrity changes with disease progression. We investigated these questions in autosomal dominant Alzheimer's disease mutation carriers, whose conserved age at dementia onset allows individual staging based upon their estimated years to symptom onset. From the Dominantly Inherited Alzheimer Network observational cohort, we selected T₁-weighted MRI scans from 269 mutation carriers and 170 non-carriers (mean age 38 ± 15 years, mean estimated years to symptom onset -9 ± 11), of whom 237 had longitudinal scans with a mean follow-up of 3.0 years. Single-subject grey matter networks were extracted, and we calculated for each individual the network properties which describe the network topology, including the size, clustering, path length and small worldness. We determined at which time point mutation carriers and non-carriers diverged for global and regional grey matter network metrics, both cross-sectionally and for rate of change over time. Based on cross-sectional data, the earliest difference was observed in normalized path length, which was decreased for mutation carriers in the precuneus area at 13 years and on a global level 12 years before estimated symptom onset. Based on longitudinal data, we found the earliest difference between groups on a global level 6 years before symptom onset, with a greater rate of decline of network size for mutation carriers. We further compared grey matter network small worldness with established biomarkers for Alzheimer disease (i.e. amyloid accumulation, cortical thickness, brain metabolism and cognitive function). We found that greater amyloid accumulation at baseline was associated with faster decline of small worldness over time, and decline in grey matter network measures over time was accompanied by decline in brain metabolism, cortical thinning and cognitive decline. In summary, network measures decline in autosomal dominant Alzheimer's disease, which is alike sporadic Alzheimer's disease, and the properties show decline over time prior to estimated symptom onset. These data suggest that single-subject networks properties obtained from structural MRI scans form an additional non-invasive tool for understanding the substrate of cognitive decline and measuring progression from preclinical to severe clinical stages of Alzheimer's disease.

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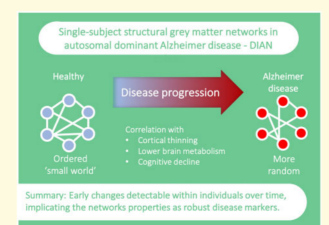
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Abbreviations: $A\beta$ = amyloid β ; ADAD = autosomal dominant Alzheimer's disease; *APP* = *amyloid precursor protein gene*; CDR = Clinical Dementia Rating scale; DIAN = Dominantly Inherited Alzheimer Network; EYO = estimated years to symptom onset; FDG = 18F-Fluorodeoxyglucose; *PSEN* = *Presenilin gene*; ROI = region of interest; SPM = Statistical Parametric Mapping software; SD = standard deviation; SUVR = standardized uptake value ratio.

Graphical Abstract



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Introduction

In order to advance clinical trials to slow or halt Alzheimer's disease, the most frequent cause of dementia (Scheltens *et al.*, 2016), it is important both to understand the evolution of pathophysiological changes occurring and to develop disease progression markers (Aisen *et al.*, 2017). Current biomarkers reliably detect Alzheimer's disease pathology (Jack *et al.*, 2018), however, predicting and monitoring disease progression remain difficult. Brain network properties are linked to cognitive function (Bassett and Bullmore, 2009; Chhatwal *et al.*, 2018; Franzmeier *et al.*, 2018), therefore studying network integrity may offer new insights into disease progression in Alzheimer's disease.

One way to measure of brain networks is by determining the similarity of grey matter morphological measures between brain regions across individuals, i.e. grey matter covariance networks (He et al., 2008; Li et al., 2012; Tijms et al., 2012) (Fig. 1). This approach is based on the notion that brain regions involved in distinct cognitive functions tend to develop in a similar way, possibly due to shared neurotrophic factors (Zielinski et al., 2010; Alexander-Bloch et al., 2013a, b). Common developmental trajectories and functional co-activation result in similar grey matter tissue properties, as measured on structural MRI (Draganski et al., 2004; Mechelli et al., 2005; Seeley et al., 2009). These covariance patterns are related to normal cognition (Seidlitz et al., 2018; Doucet et al., 2019) and reveal in healthy individuals an optimal, 'small-world', organization by graph theory description (He et al., 2007; Humphries and Gurney, 2008). In sporadic Alzheimer's disease dementia, grey matter networks are disrupted, the properties show a less optimal, random organization of the network (Yao et al., 2010; Tijms et al., 2013a; Kim et al., 2016). In pre-dementia stages, such loss of network integrity predicts clinical progression and cognitive decline (Dicks et al., 2018; Tijms et al., 2018). The presence of amyloid β (A β) pathology in cognitively normal individuals has also been associated with grey matter network alterations (Tijms et al., 2016; Ten Kate et al., 2018; Voevodskaya et al., 2018). Together, these observations suggest that these network properties change over the course of Alzheimer's disease, from early stages, and that individual grey matter network extractions could possibly be used to monitor disease progression. However, as previous findings were based on onetime network extractions, it remains unclear whether, and when, these networks change *within* individuals as they progress in their disease.

A complication when studying sporadic Alzheimer's disease is the difficulty of placing pre-symptomatic individuals on their disease timeline (Villemagne *et al.*, 2013; Donohue *et al.*, 2014; Young *et al.*, 2014; Roe *et al.*, 2018; Vermunt *et al.*, 2019). This issue is less problematic for carriers of a genetic mutation that causes autosomal dominant Alzheimer's disease (ADAD), because the age at onset of dementia can be estimated, from the age at onset in family members or carriers of the same specific mutation type. The estimated years to symptom onset (EYO) can serve as a proxy for disease duration (Bateman *et al.*, 2012; Ryman *et al.*, 2014). Using this paradigm, previous work demonstrated that $A\beta$ aggregation starts more than two decades before dementia onset (Gordon *et al.*, 2018; McDade *et al.*, 2018; Oxtoby *et al.*, 2018). Closer to symptom onset, individuals show accelerated hypometabolism and cortical thinning, which is followed by cognitive decline (Benzinger *et al.*, 2013; Kinnunen *et al.*, 2018; Wang *et al.*, 2019). When during these processes, grey matter networks start to decline remains unknown.

Here, we investigated for the first time single-subject grey matter networks over the course of ADAD. We assessed when, and how, the network properties change as a function of EYO, both cross-sectionally and longitudinally, on a global and regional level. To understand the relationship between grey matter network property changes and disease progression, we also investigated how the network smallworld coefficient alters with established Alzheimer's disease markers of $A\beta$ accumulation, brain metabolism, cortical thickness and cognitive function.

Materials and methods

DIAN study design and participants

In the worldwide Dominantly Inherent Alzheimer Network (DIAN) longitudinal cohort study, families with individuals carrying a PSEN1, PSEN2 or APP mutation undergo genetic testing and repeated clinical, cognitive, fluid and brain imaging assessments. The non-carrier family members act as an inherent control group. Participants generally have study visits every 3 years at earlier disease stages and these assessments become yearly when either symptoms are present, or they are within 3 years of their EYO. DIAN protocols had supervisory approval from the ethical review board of Washington University in St. Louis, and all participants gave informed consent. For this study, we selected data from all participants who had undergone at least one MRI scan that passed quality control in the 12th data freeze. Families with the Dutch or Flemish APP mutation were excluded because these mutations result in a different phenotype, with predominantly cerebral amyloid angiopathy.

Estimated years to symptom onset

We calculated the EYO for mutation carriers and noncarriers identically: The EYO was defined as the mutation-specific mean age at onset subtracted by the individuals' visit age (Ryman *et al.*, 2014). In case of an unknown mutation-specific age at onset, the parental age at disease onset, reported by the participant, was used

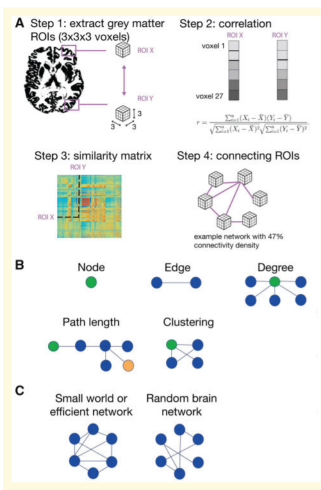


Figure | Details on grey matter network metrics. (With permission from Verfaillie et al. 2018, Human Brain Mapping.) (A) Grey matter network extraction from the individual brain segmentation (described in text). (B) The sum of the number of nodes, i.e. the number of cubes, is the size of the network. The degree is the average number of connections per node. The connectivity density is the percentage of the number of connections in the network compared to the maximum number of connections possible. The clustering coefficient of a node describes the proportion of connections between neighbours for every node. For example, in case a node connects to 3 other nodes, there are 3 possible connections between those 3 adjacent nodes. If only I connection is present between 2 of the 3 other nodes, the clustering of the primary node is 1 out of 3, 0.33. Global clustering is determined by taking averaging clustering values across all nodes. Path length is the mean of the shortest paths for a node to reach every other node in the network. The global path length is the average path length across all nodes. (C) Normalized clustering and normalized path length describe how on a global level a network organization differs from a randomly organized network. The networks are randomized by rewiring the connections randomly in each network, while keeping intact the total number of nodes and degrees (Maslov and Sneppen, 2002). The network's observed clustering and path length are divided by the clustering and path length values, respectively, of averaged random networks to obtain the normalized values. Lastly, the small-world coefficient is the normalized clustering divided by the normalized path length. The network has the 'small-world property' if this ratio is higher than I, indicating a path length close to the random networks, yet a greater then random clustering. This is optimal, because of fast exchange of information between remote clusters, and specialized information processing within clusters.

instead. For example, if the mean age at symptom onset for a specific mutation is 50 years, then a 35-year-old individual would have an EYO of -15. For the carriers of the ADAD mutation, this indicates that the individual is expected to show clinical symptoms of Alzheimer's disease 15 years later.

Clinical evaluation and cognition

Disease severity was measured using the Clinical Dementia Rating scale (CDR) (Morris, 1993), administered to the participant and study partner by blinded raters. Participants were classified as being unimpaired (global CDR score = 0) or symptomatic (global CDR 0.5, 1, 2 and 3). In addition, cognitive function was summarized using a cognitive composite developed in the DIAN project (Bateman *et al.*, 2017), consisting of the average of equally weighted Z-scores of the Logical Memory delayed recall total score from the Wechsler Memory Scale-Revised, DIAN Word List Test delayed free recall score, Digit Symbol Coding total score from the Wechsler Adult Intelligence Scale-Revised Digit Symbol Substitution Test, and the total score from the Mini-Mental State Examination.

MRI acquisition and pre-processing

MRI T₁-weighted scans $(1.1 \times 1.1 \times 1.2 \text{ mm}^3 \text{ voxels}, \text{repetition time} = 2300 \text{ ms}, \text{ echo time} = 2.95 \text{ ms}, \text{ flip angle 9°}) were acquired according to Alzheimer's Disease Neuroimaging Initiative (ADNI) protocols (Jack$ *et al.* $, 2010). We segmented T₁ images into grey and white matter and CSF, using the Statistical Parametric Mapping software version 12 (SPM12; Wellcome Trust Centre for Neuroimaging, UCL Institute of Neurology, London, UK). All segmentations were checked visually, after which 51 scans were removed due to failed segmentations or severe motion artefacts. Native space grey matter segmentations were resampled into <math>2 \times 2 \times 2 \text{ mm}^3$ voxels. This voxelwise data were used as input for connectivity analyses.

Single-subject grey matter networks and metrics

Grey matter networks were computed according to a previously published, automated pipeline (Tijms *et al.*, 2012) that includes two steps figurated in Fig. 1A: (i) grey matter network extraction (https://github.com/bettytijms/Single_ Subject_Grey_Matter_Networks, accessed December 2019; implemented in Matlab2016b (MathWorks, Natick, MA)) and (ii) graph theory-based metric calculation (Rubinov and Sporns, 2010; Tijms *et al.*, 2012). To extract singlesubject grey matter networks, we parcellated each individual's native space grey matter segmentation into $6 \times 6 \times 6$ mm³ cubes, containing 27 voxels. These non-overlapping cubes serve as the 'nodes' in the network. Connections between each pair of cubes across an individual's scan were established by calculating the Pearson's correlation coefficient between the corresponding voxels. This approach takes into account both the grey matter probability (i.e. from the tissue segmentation) as well as the spatial information present in 27 voxels within each cube. All correlations were stored in a matrix, and the presence or absence of connections between nodes was dichotomized according to an individualized threshold that ensured a maximum of 5% spurious connections for each individual (Tijms *et al.*, 2012).

For each individual's binarized grey matter network, we calculated graph theory metrics describing the global network properties: size, degree, connectivity density, clustering coefficient, path length, normalized clustering, normalized path length and small-world coefficient (see Fig. 1B and C for explanation of these metrics). We also calculated regional network properties. In order to aid comparability with other studies previously performed in DIAN, regional network metrics were calculated within each region of the Desikan-Killiany atlas (Desikan et al., 2006). The regional masks were obtained by first parcellating each individual's T1 image into 34 anatomical regions of interest (ROIs) from the Desikan atlas using Freesurfer 5.3 (Fischl, 2012) (http://surfer.nmr.mgh.har vard.edu, accessed December 2019). The Freesurfer output was then aligned to the native space T_1 using FSL (https://fsl.fmrib.ox.ac.uk/fsl, accessed December 2019), and this transform was used to register the parcellation into native space. The network values of the degree, clustering coefficient and path length were subsequently averaged within a region. Graph theory metrics were calculated using scripts from the brain connectivity toolbox (https://sites.google.com/site/bctnet/, accessed December 2019), modified for large-sized networks.

Other DIAN imaging data

We examined regional data for $A\beta$ using PET imaging with 11 C-Pittsburgh Compound B ($A\beta$ PET), glucose metabolism with 18F-Fluorodeoxyglucose PET (FDG-PET), and cortical thickness and volumes from structural MRI. Details on data processing have previously been described (Gordon *et al.*, 2018). The Freesurfer ROIs were used to process the amyloid and FDG-PET data. PET data are processed using a cerebellar grey reference region and partial volume corrected using a geometric transfer matrix approach (Su *et al.*, 2013; Su *et al.*, 2015). In this study, we utilized the MRI precuneus cortical thickness, the precuneus $A\beta$ PET and to match a previously defined meta-ROI, the average of the left and right isthmus cingulate and inferior parietal region in FDG-PET for crossmodal comparison with grey matter network properties (Landau *et al.*, 2011).

Statistical analyses

As part of sample characterization, we compared four groups (non-carriers; asymptomatic mutation carriers with an EYO before -15 years; asymptomatic mutation carriers with EYO between -15 and 0 years; and symptomatic mutation carriers) on cross-sectional grey matter network small-world values, and the other network measures with the Kruskal–Wallis test, and *post hoc* Wilcoxon test with Holm *P*-value adjustment. We extracted individual slopes with linear mixed models in R for those individuals with repeated measures. Using the same statistical tests, we compared those extracted slopes between the groups. In addition, we compared individuals with different mutation types (PSEN1/PSEN2/APP) on the baseline network property values.

For the main analyses, we compared mutation carriers and non-carriers to determine (i) the EYO at which grey matter network metrics showed cross-sectional differences between groups and (ii) the EYO at which the groups had a different rate of change over time by fitting linear mixed effects models. Specifically, we used Bayesian inference methods (Gordon et al., 2018; Mishra et al., 2018) to determine the EYO point that 99% credible intervals of the difference distribution did not overlap 0. In these methods, the model parameters were estimated as previously described, applying a Hamiltonian Markov chain Monte Carlo sampling of the posterior distribution, with 10 000 iterations in eight chains, 5000 warm-up, thinning retaining 1 out of every 10 iterations and cauchy prior in the STAN package for R. We checked the model convergence statistic Gelman-Rubin diagnostic, the Rstatistic (rhat), which compares the between and withinchain estimates for each of the model parameters. These should be at least close to 1.0 and were for all models close to 1.00. (Gelman et al., 2015; Carpenter et al., 2017). From the posterior distribution, we calculated the range 99% credible intervals around the estimates, i.e. 0.005-0.995 range. We also calculated the difference curve between the mutation carriers and non-carriers by EYO with 99% credible intervals. We refer to the 'divergence point' as the point where the 99% credible interval of this difference curve between carriers and non-carriers did not contain 0 (i.e. no difference). The credible interval is the Bayesian equivalent of the frequentist confidence interval. The main difference is that the Bayesian directly estimates the credible interval from an actual computed population (i.e. posterior) distributions, rather than hypothesized as in the frequentists approach. Therefore, an advantage of the Bayesian approach is that the credible interval can be interpreted in a probabilistic way. To allow for non-linear effects, without assuming a particular shape, we applied a restricted cubic spline with knots at the 0.10, 0.50 and 0.90 of the EYO

distribution, also described previously (Gordon et al., 2018) that included a linear term (EYOlinear) and a cubic term (EYOcubic). Cross-sectional models contained fixed terms for EYO, mutation status, their interaction and a random effect for family cluster. Longitudinal models were used to study the rate of change of network properties, and individuals with one data point were also included. Those models included fixed terms for baseline EYO (two terms: EYOlinear and EYOcubic), time after baseline, mutation status and, all two- and three-way interactions (see formulas in Supplementary material, p. 6). Additionally, all models included random intercept terms for subject and family cluster and a random slope for subject. The covariates whole-brain grey matter volume and sex were included as fixed terms. Equivalent to previous work, when size, degree or connectivity density were found to be associated with mutation status in any of the models, were included as additional covariates for sensitivity analysis as these variables also influence more complex network metrics (Tijms et al., 2013a). Regional models were adjusted for sex, regional degree and regional grey matter volume.

We examined relationships between grev matter network small-world coefficient and established Alzheimer's disease markers within mutation carriers. Previous research suggested grey matter networks may be disrupted in response to $A\beta$ accumulation, precipitating cognitive decline (Ten Kate et al., 2018). For this reason, our models included either precuneus PET A β as a predictor and grey matter network metrics as outcomes or grey matter network metrics as a predictor and cortical thickness (precuneus), brain metabolism (meta-ROI), or cognition (DIAN cognitive composite) as the respective outcomes. These predictors and outcomes were Z-scored to the whole group. We fitted three sets of linear mixed effects models that were all adjusted for baseline grey matter volume, age, and sex, and with random intercept for family cluster, in lme4 package in R (Bates et al., 2014) (see detailed formulas in Supplementary material, p. 6). If models failed to converge, the term for family cluster was removed. Models were divided into three sections. The first was baseline comparisons. The second set was longitudinal comparisons in participants with at least two data points to avoid convergence issues and included additional random effects for subject intercept and slope of the predictor. The final set of models was used to evaluate whether baseline data could predict change over time in the outcome. These models had fixed effects for baseline predictor, time from baseline, and its interaction, and a random subject intercept and slope of time from baseline. We focused on the grey matter network smallworld coefficient, as this metric is indirectly derived from all other network metrics, and can thus be considered a summary statistic (Fig. 1). We also show exploratory graphs for the other network measures for completeness and repeated as a sensitivity analysis the cross-modal models for the mutation carriers only.

Data availability

The data from the DIAN study can be requested online at https://dian.wustl.edu/, accessed December 2019.

Results

In total, 439 participants from the DIAN study, with a mean \pm SD age of 38 \pm 11 years and a mean \pm SD EYO of -9 ± 11 , had MRI scans of sufficient quality to be included in the present analyses. The group consisted of 269 (61%) ADAD mutation carriers and 170 (39%) non-carrier family members (Table 1). Of this sample, 237 (54%) participants had longitudinal MRI scans, with a mean of 2.5 scans per participant and a maximum of 6 acquired over a mean ± SD 3.0 ± 1.5 years of follow-up (clinical and PET data in Supplementary Table 1). There were groups differences between asymptomatic mutation carriers with EYO < -15 years, asymptomatic mutation carriers -15 < EYO < 0 years, symptomatic mutation carriers and non-carriers on cross-sectional network values and extracted slopes (Supplementary Figs 3 and 4). For most network measures, we found that the mutation carriers who are far away from expected onset (EYO >15 years) and the non-carriers had slightly higher network property value than mutation carriers who were closer to expected symptom onset, and it further decreased in the symptomatic stage. Rate of decline showed a similar pattern between these groups. Figure 2 illustrates these comparisons for the small-world coefficient. Comparing PSEN1/PSEN2/APP mutation carriers at baseline on all network metrics, the network size and average degree were slightly lower in PSEN1 carriers, while the other metrics were similar (Supplementary Fig. 5).

Cross-sectional divergences between mutation carriers and non-carriers

The mutation carriers diverged from non-carriers on all grey matter network metrics, except for network size and raw path length (Fig. 3, Supplementary Table 2). Lower network metric values for mutation carriers relative to non-carriers were observed earliest in normalized path length at EYO -12, followed by lower normalized clustering at EYO -8.7, small-world coefficient at EYO -8.4, clustering coefficient at EYO -7.5, connectivity density at EYO -5.6 and degree at EYO 0. When additionally adjusting for degree or connectivity density, the estimates for network metrics yielded similar results (Supplementary Table 2). Using the same methods, but now implemented on a regional level, the earliest divergence between mutation carriers relative to non-carriers was found for path length in the precuneus at EYO -13.1, for clustering in the superior temporal gyrus at EYO -10 and for network degree in the banks of the superior temporal gyrus at EYO -7 (Fig. 4, Supplementary Table 3).

Table 3).

and cognition

Table | Group characteristics

Divergences of rates of change

between mutation carriers

Association of grey matter

networks with other neuroimaging

Established markers of Alzheimer's disease showed signifi-

cant relationships with the small-world coefficient used as

a global network summary statistic. We examined cross-

modal relationships between baseline markers; over

repeated measures; and whether baseline values could predict further decline in the other marker. We found

that higher A β deposition load on PET was cross-sectionally related to a lower small-world coefficient ($\beta \pm SE =$

 -0.22 ± 0.05 , $P = 3 \times 10^{-6}$, Fig. 5). In a longitudinal de-

sign, faster amyloid accumulation over time related to

concurrent small-world coefficient decline ($\beta \pm SE =$

 -0.33 ± 0.06 , $P = 1 \times 10^{-7}$). Thirdly, a higher amyloid

load at baseline predicted steeper decline of the small-

compared to non-carriers

world coefficient over time ($\beta \pm SE = -0.07 \pm 0.01$, $P = 4 \times 10^{-8}$).

Grev matter network small-world coefficient and the markers of Alzheimer's disease progression showed sig-When comparing rates of change over time, mutation nificant relationships, both cross-sectionally and longitucarriers diverged from non-carriers by EYO for all grey dinally (Fig. 6). Specifically, a lower small-world matter network metrics, except connectivity density. coefficient was cross-sectionally related to lower FDG-Steeper decline for mutation carriers relative to non-car-PET metabolism in the meta-ROI ($\beta \pm SE = 0.44 \pm 0.08$, riers was detected earliest for network size, at baseline $P=2 \times 10^{-7}$), as well as lower precuneus cortical thick-EYO -6.0, followed by small-world coefficient at EYO ness ($\beta \pm SE = 0.50 \pm 0.06$, $P = 2 \times 10^{-15}$). For cogni--4.7, normalized clustering at EYO -4.6, degree at EYO tion, a lower small-world coefficient was cross-sectionally -4.4, normalized path length at EYO -2.8, clustering related to lower scores on the DIAN cognitive composite coefficient at EYO -2.6 and path length at +1.0 (Fig. 2, $(\beta \pm SE = 0.28 \pm 0.08, P = 3 \times 10^{-4})$. In a longitudinal Supplementary Table 2 and Figs 1 and 2). When adddesign, decline of the small-world coefficient over time itionally adjusting for degree, the estimates for network related to concurrent decreases of FDG-PET metabolism metrics yielded similar results, except for clustering coeffi- $(\beta \pm SE = 0.54 \pm 0.06, P = 5 \times 10^{-14})$ and faster precucient, which lost significance. On a regional level, the earneus cortical thinning ($\beta \pm SE = 0.55 \pm 0.06$, $P = 1 \times$ liest steep decline rate for mutation carriers compared to 10^{-17}). A declining small-world coefficient over time was non-carriers was detected for degree in the lateral occipirelated to concurrent decline on the cognitive composite tal gyrus at EYO -7.4, for clustering in the parahippo- $(\beta \pm SE = 0.47 \pm 0.06, P = 2 \times 10^{-11})$. Thirdly, a lower campal gyrus at EYO -6.2 and for path length in the small-world coefficient at baseline predicted faster neuroprecentral gyrus at EYO -4.2. (Fig. 4, Supplementary degeneration as measured by FDG-PET metabolism ($\beta \pm$ SE = 0.12 ± 0.02 , $P = 2 \times 10^{-8}$) and precuneus cortical thinning $(\beta \pm SE = 0.10 \pm 0.01, P = 4 \times 10^{-12})$, and steeper cognitive decline over time (composite $\beta \pm SE =$ 0.08 ± 0.02 , $P = 2 \times 10^{-7}$). Associations for the other network properties can be found in Supplementary Figs 6–9 in explorative graphs.

> We repeated the cross-modal analyses, this time solely including the mutation carriers who were asymptomatic at baseline (see Supplementary Table 4). In brief, most relationships, albeit attenuated, were also present in the asymptomatic mutation carriers only. The cross-sectional relationships with the small-world coefficient remained significant for FDG-PET metabolism and precuneus cortical thickness. All longitudinal relationships indicating concurrent changes between markers were significant. Of the third set of models, aimed at predicting the change over time, two models did not converge (with amvloid PET and with cognition). The baseline small-world property still predicted the decline of FDG-PET metabolism.

	Non-carriers (N = 170)	Asymptomatic mutation car- riers (N = 174)	Symptomatic mutation car- riers (N = 95)
Baseline age, years	38 (11)	34 (9)	46 (10)
Female, N (%)	101 (59%)	100 (57%)	50 (53%)
Estimated years to onset	-11(12)	-14 (8)	I (7)
MMSE	29.1 (1.2)	29.1 (1.2)	22.9 (6.6)
Total MRI scans, 1/2/3/4–6, N	84/61/18/7	84/59/28/3	34/30/17/14
Longitudinal scans, mean (SD)	2.4 (0.8)	2.4 (0.7)	2.9 (1.1)
Follow-up time MRI visits, years	3.3 (1.5)	3.2 (1.5)	2.2 (1.3)
Mutation type PSEN1/PSEN2/APP N	n/a	133/16/25	75/2/18

Mean (SD), unless otherwise specified. EYO is the expected age at onset of the mutation that runs in the family. MMSE = Mini-Mental State Examination.

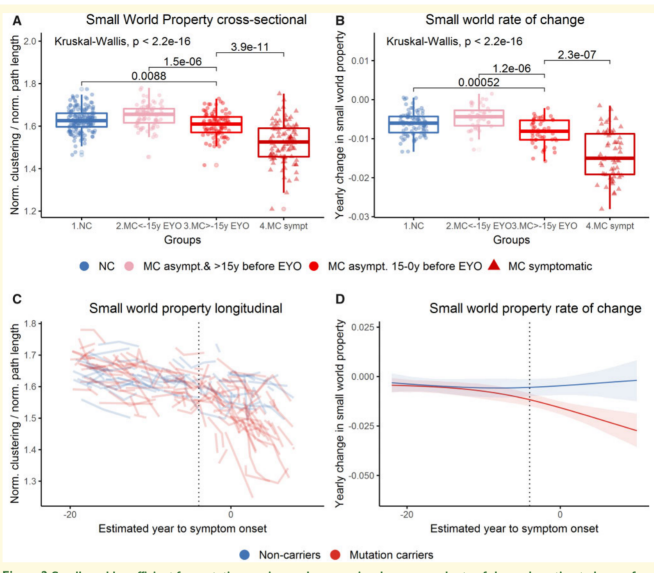


Figure 2 Small-world coefficient for mutation carriers and non-carriers by group and rate of change by estimated year of onset. (A, B) *P*-values based on Kruskal–Wallis, and *post hoc* Wilcoxin with Holm method correction shown for the comparison of all groups to the asymptomatic EYO between -15 and 0 mutation carriers group. No covariates included. (C, D) The fitted lines are based on all data points extending to -38 to +20. Left of EYO 0 is before expected symptom onset and right of EYO 0 is after expected symptom onset. The EYO were first jittered, and then the data points before -20 and after EYO +8 removed to avoid accidental unblinding of participants. Dotted line is the point of divergence of rate of change between mutation carriers and non-carriers.

Discussion

Using a single-subject approach, we found that structural grey matter network properties deteriorated over the course of ADAD and that movement to a more random network topology closely correlated with cognitive decline. When comparing mutation carriers to non-affected family members global network disruptions were detected cross-sectionally as early as 12 years before expected symptom onset. Longitudinally, increased rates of decline of network metrics were evident from 6 years before expected symptom onset. In line with our hypotheses based on cross-sectional studies in sporadic Alzheimer's disease, a lower small worldness of the network was associated with abnormalities and decline of established markers of Alzheimer's disease. Thus, our grey matter network analysis in this unique cohort of ADAD contributes to our understanding of the Alzheimer's disease trajectory and indicates that our methods may potentially be a useful additional non-invasive tool for tracking disease progression.

As Alzheimer's disease progresses, there is substantial amyloid accumulation, volumetric loss, hypometabolism and cognitive decline, but how grey matter networks fit into these processes remained unclear. Prior work in sporadic Alzheimer disease has shown that grey matter networks might be sensitive to biological changes during the preclinical stages of the disease (Tijms *et al.*, 2016;

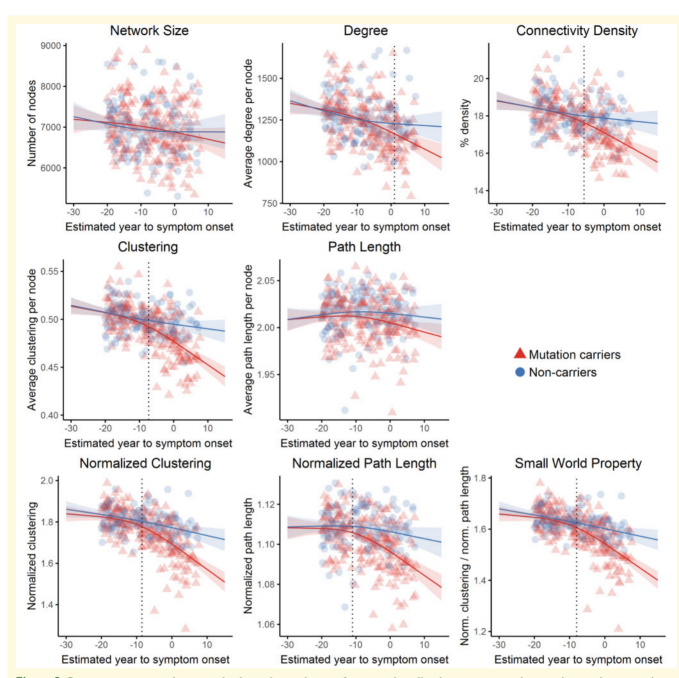


Figure 3 Grey matter network properties by estimated year of onset at baseline between mutation carriers and non-carriers. The fitted lines are based on all data points extending to -38 to +20. Left of EYO 0 is before expected symptom onset and right of EYO 0 is after expected symptom onset. The EYO were first jittered and then the data points before -20 and after EYO +8 removed to avoid accidental unblinding of participants. Dotted line is the point of divergence between mutation carriers and non-carriers. N = 439.

Ten Kate *et al.*, 2018; Voevodskaya *et al.*, 2018). In the current work, we observed similar alterations of grey matter network properties in ADAD as a function of EYO. The mostly consistent changes in network properties between sporadic and ADAD strengthens the hypothesis that grey matter network disruptions are one of the downstream effects of amyloid accumulation. Using amyloid PET, we extended previous cross-sectional findings from studies in sporadic Alzheimer's disease (Ten Kate

et al., 2018), by showing that higher baseline amyloid PET and higher amyloid accumulation rates are related to faster decline of grey matter network properties over time. Within asymptomatic mutation carriers only, the relationship between amyloid and the small-world coefficient was more subtle and only reached significance when studying concurrent changes of both markers, possibly due a decrease in power. The small-world summery measure was also related to sensitive markers of

Alzheimer's disease neurodegeneration and cognitive decline, in cross-sectional and longitudinal design. For these relationships, the sensitivity analyses in asymptomatic mutation carriers showed that the small-world coefficient already in early disease stages declined concurrently with other Alzheimer's disease markers. This suggested these processes occur, at least partly, in parallel (Wang *et al.*, 2019), and support the notion that grey matter network decline is a sign of progression of Alzheimer's disease.

Previous studies in sporadic Alzheimer's disease had suggested decline over time of grey matter network integrity, as there was a decrease over disease stages cross-sectionally (Yao *et al.*, 2010; Tijms *et al.*, 2013b; Voevodskaya *et al.*, 2018). Here, we show that grey matter networks properties decline over time *within* individuals, and how decline rates start to increase with disease severity. Differences between mutation carriers and non-

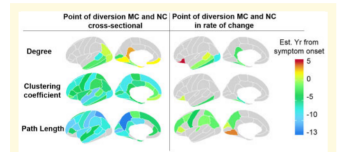


Figure 4 Regional EYO of diversion between mutation carriers and non-carriers for grey matter network degree, clustering coefficient and path length. Linear mixed models adjusted for sex, total grey matter volume and regional volume. MC = mutation carrier; NC = non-carrier. For details, EYO by region see Supplementary Table 3. N = 416.

carriers in the *rate* of decline were generally detected later than cross-sectionally, which could have occurred because cross-sectional estimates across individuals by EYO may overestimate changes due to variance in the EYO measure (i.e. some individuals at EYO -12 are actually only 5 or 6 years from actual onset) (McDade et al., 2018). Another potential cause of cross-sectional and longitudinal estimate differences include sample sizes, with less indiwho had longitudinal data. Measurement viduals variability over repeated measures within individuals can also have contributed to later detection of differences in the longitudinal design if these exceeded subtle rates of change. Longer follow-up time per individual in larger visit numbers is necessary for increasingly precise estimates of divergence in change over time.

Altering of network properties was not detected for every metric. This may be an indication that these metrics pick up different aspects of neurodegeneration. The small-world measures (normalized clustering, and normalized path length and small-world coefficient) showed early cross-sectional changes and seemed most sensitive to measure change over time. This is in line with network theory and previous findings in Alzheimer's disease (Tijms et al., 2018), which indicated that brain networks tend to become more similar to random networks over the disease course. The normalized network metrics reflect how different a network is from random, which may be why these best capture decline over time. Future studies are needed to confirm which network property would be the most robust summary statistic to track longitudinal grey matter network integrity in Alzheimer's disease.

On a regional level, cross-sectional network property alterations were evident earliest in the parietal regions and then spread across the brain. Most brain regions

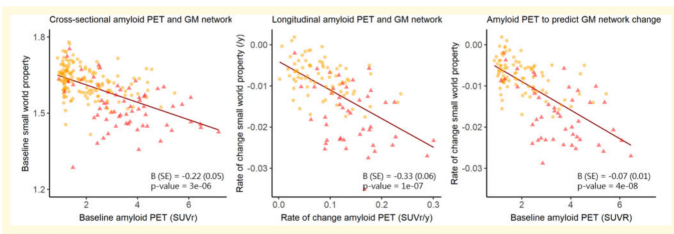
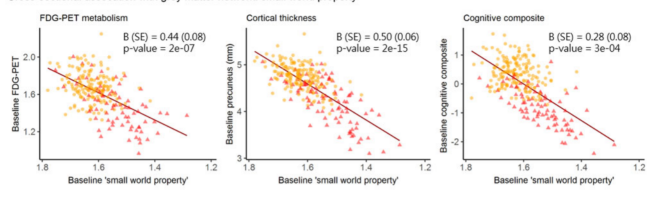
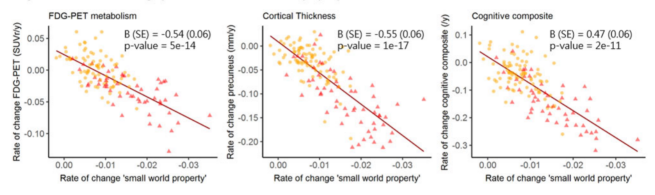


Figure 5 Association of amyloid PET with grey matter network small-world coefficient in mutation carriers. For visualization purposes, plotted extracted slopes with mixed model and line fitted with simple regression line in ggplot in R. Models to obtain beta and *P*-values specified in methods. GM network = grey matter network. Yellow circle = CDR 0 at baseline; Red triangle = CDR >0 at baseline. Amyloid PET = precuneus SUVr, Cross-sectional N= 222, Longitudinal N = 120, Predict change N = 131. For other grey matter network metrics see Supplementary Fig. 6.

Cross-sectional assocation with grey matter network: small world property



Longitudinal assocation with grey matter network: small world property



Predict change with baseline grey matter network: small world property

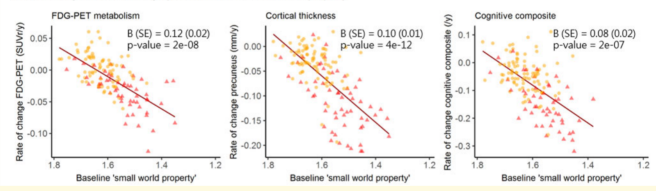


Figure 6 Associations of grey matter network small-world coefficient with FDG-PET metabolism, cortical thickness and cognition. For visualization purposes, plotted extracted slopes with mixed model and line fitted with simple regression line in ggplot in R. Models to obtain beta and *P*-values specified in methods. Inversed small-world coefficient to aid visualization, see also Supplementary Table 4. Yellow circle = CDR 0 at baseline; Red triangle = CDR >0 at baseline. MRI thickness = cortical thickness precuneus; FDG-PET = METAROI SUVr as described in methods. DIAN composite: equally weighted *Z*-score of Logical Memory Delayed Recall of the Wechsler memory test, DIAN Word List Test (comparable to International Shopping List Test), Digit Symbol Substitution Test and Mini-Mental State Examination. Cross-sectional FDG-PET N = 238, MR thickness N = 260, Cognition N = 251; Longitudinal: FDG-PET N = 129, MR thickness N = 146, Cognition N = 143. For other grey matter network metrics see Supplementary Figs 7–9.

showed a difference first for path length, then for clustering and then for degree, except for the temporal regions, in which earlier and more pronounced lowering of the clustering coefficient was seen. Regional cross-sectional patterns showed early alterations for path length and clustering in areas with most pathology in ADAD including the precuneus. Regions of the default mode network also showed early alterations. Compared to previous sporadic Alzheimer's disease studies, we find more widely affected connectivity but the patterns are largely overlapping (Ten Kate *et al.*, 2018; Tijms *et al.*, 2018; Verfaillie *et al.*, 2018).

Compared to other structural grey matter imaging, the cross-sectional differences in the most sensitive grey matter network metrics were detected earlier than cortical thickness and volumetric measures. It was not part of this study to investigate whether grey matter network integrity measures have the same or higher sensitivity to early alterations than other structural MRI markers. Still, we adjusted for grey matter volume to assure measuring value beyond simple volumes. The increased rates of change of network properties were detected at a similar stage to the volumetrics, and later than precuneus cortical thinning in dominantly inherited Alzheimer's disease, which is the earliest region of change (Gordon et al., 2018; Kinnunen et al., 2018). The results merit application of grey matter networks in future deeper investigations, for example using multimodal network approaches with white matter and functional connectivity, to better understand the substrate of cognitive decline. The observation that network disruptions increase over time in a large multicentre study is relevant for clinical trials. As the method only requires standard T1 scans and the available pipeline for network calculation, a next step is to test the approach retrospectively in clinical trial populations.

One of the strengths of the current study design is the use of a previously validated method for grey matter network extraction. The unique traits of the DIAN cohort provided the ability to map changes in grey matter networks across decades of disease time. It should be noted that the estimates as a function of the expected symptom onset in dominantly inherited Alzheimer's disease are influenced by sample size. Still, this method provides a way to detect and compare changes due to Alzheimer's disease before symptom onset, and combine different families. Additionally, the rich characterization of DIAN participants provided the ability to relate observed changes in networks to other neuroimaging markers of pathology as well as cognition. A potential limitation is that our study included an average time period of 3 years in the longitudinal cohort, which may not be enough time to reliably measure changes due to Alzheimer's disease in its very early stages. Yet, we show the longitudinal analysis of structural grey matter networks alongside of the cross-sectional results, which to the best of our knowledge has not been studied before and warrants further investigation of how grey matter network integrity decreases over time in sporadic Alzheimer's disease.

In conclusion, in ADAD individual grey matter network properties are robustly associated with Alzheimer's disease severity and progression as shown by the associations with EYO, amyloid accumulation, rate of neurodegeneration and cognitive decline. These data suggest that single-subject grey matter network integrity measures obtained from structural MRI scans provide an additional, non-invasive tool for understanding and measuring progression from preclinical to severe clinical stages of Alzheimer's disease. These grey matter network properties can reflect the asynchronous start of brain pathology following Alzheimer' disease-related cellular damage and inflammatory processes, informing about changes in grey matter covariance (Verfaillie *et al.*, 2018).

Supplementary material

Supplementary material is available at Brain Communications online.

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Competing interests

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Appendix I. The dominantly inherited Alzheimer's network

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