A longitudinal study on resting state functional connectivity in behavioral variant frontotemporal dementia and Alzheimer’s disease

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Abbreviations

AD: Alzheimer’s disease
bvFTD: behavioral variant of Frontotemporal Dementia
CDR: Clinical Dementia Rating Scale
FAB: FMRIB’s ICA-based Xnoiseifier
fMRI: Functional Magnetic Resonance Imaging
FSL: Functional Magnetic Resonance Imaging of the Brain Software Library
FWE: Family-Wise Error
GDS: Geriatric Depression Scale
GLM: General Linear Model
ICA: Independent Component Analysis
MMSE: Mini-Mental State Examination
MNI: Montreal Neurological Institute
MRI: Magnetic Resonance Imaging
TFCE: Threshold-Free Cluster Enhancement
VBM: Voxel-Based Morphometry
Abstract

Background and objective: Alzheimer’s disease (AD) and behavioral variant frontotemporal dementia (bvFTD) are the most common types of early-onset dementia. We applied longitudinal resting state fMRI to delineate functional brain connections relevant for disease progression and diagnostic accuracy.

Methods: We used two-center resting state fMRI data of 20 AD patients (65.1±8.0 years), 12 bvFTD patients (64.7±5.4 years), and 22 control subjects (63.8±5.0 years) at baseline and 1.8-year follow-up. We used whole-network and voxel-based network-to-region analyses to study group differences in functional connectivity at baseline and follow-up, and longitudinal changes in connectivity within and between groups.

Results: At baseline, connectivity between paracingulate gyrus and executive control network, between cuneal cortex and medial visual network, and between paracingulate gyrus and salience network was higher in AD compared with controls. These differences were also present after 1.8 years. At follow-up, connectivity between angular gyrus and right frontoparietal network, and between paracingulate gyrus and default mode network was lower in bvFTD compared with controls, and lower compared with AD between anterior cingulate gyrus and executive control network, and between lateral occipital cortex and medial visual network. Over time, connectivity decreased in AD between precuneus and right frontoparietal network and in bvFTD between inferior frontal gyrus and left frontoparietal network. Longitudinal changes in connectivity between supramarginal gyrus and right frontoparietal network differ between both patient groups and controls.

Conclusion: We found disease-specific brain regions with longitudinal connectivity changes. This suggests the potential of longitudinal resting state fMRI to delineate regions relevant for disease progression and for diagnostic accuracy, although no group differences in longitudinal changes in the direct comparison of AD and bvFTD were found.
Introduction

Alzheimer’s disease (AD) and behavioral variant frontotemporal dementia (bvFTD) are the most common types of early-onset dementia [1]. AD is mainly characterized by deficits in episodic and working memory [2], whereas patients with bvFTD typically present with changes in behavior [3].

A substantial amount of dementia research used neuroimaging to elucidate the pathophysiology of bvFTD and AD [4,5]. Imaging of brain structure shows typical AD atrophy in the hippocampus, precuneus, posterior cingulate cortex, parietal, and occipital brain regions [6–8]. In bvFTD, atrophy is most often found in the anterior cingulate cortex, frontoinsula, and frontal brain regions [7,8].

Longitudinal studies have shown to be useful to elucidate changes in gray matter volume over time, showing that in AD atrophy progresses faster in the hippocampus and posterior cingulate cortex, while atrophy progresses faster in the orbitofrontal gyrus and frontal lobe in bvFTD [7,9–11]. These studies show the importance of longitudinal designs to delineate regions relevant for disease progression.

The disease-specific patterns of gray matter atrophy show spatial overlap with functional brain networks [8]. Imaging of these functional networks offers the opportunity to study brain function and dysfunction in AD and bvFTD [8,12]. AD patients show abnormalities in functional network connectivity in the posterior hippocampal-cingulo-temporal-parietal network known as the default mode network [13–16]. Patients with bvFTD show functional connectivity abnormalities in the anterior frontoinsular-cingulo-orbitofrontal network often called the salience network [16–21].

Despite evidence from cross-sectional studies that functional network connectivity gives the opportunity to study brain dysfunction in dementia and therefore has potential to study disease progression, little is known about how functional connections change over time in AD and bvFTD. Studying longitudinal functional connectivity is important to monitor disease progression and may
have utility to improve differential diagnosis of both types of dementia. The aim of the present study was to delineate functional connections relevant for disease progression and diagnostic accuracy. We used longitudinal resting state functional magnetic imaging (fMRI) data of AD and bvFTD patients to investigate, in addition to cross-sectional group differences, longitudinal changes in functional connectivity within and between groups.
Materials and methods

Participants
We included 20 patients with probable AD, 12 patients with probable bvFTD, and 22 control participants (Table 1). All subjects were recruited from two Dutch centers specialized in dementia: the Alzheimer Center of the VU University Medical Center Amsterdam, and the Alzheimer Center of the Erasmus University Medical Center Rotterdam.

All patients underwent a standardized dementia screening including medical history, informant-based history, physical and neurological examination, blood tests, extensive neuropsychological assessment, and magnetic resonance imaging (MRI) of the brain. In total 5 patients were treated with anticholinesterase medication and 5 patients with psychotropic medication (anxiolytics or antidepressants). Diagnoses were established in a multidisciplinary consensus meeting according to the core clinical criteria of the National Institute on Aging and the Alzheimer’s Association workgroup for probable AD [2,22] and according to the clinical diagnostic criteria for bvFTD [3]. To minimize center effects, all diagnoses were re-evaluated in a panel including clinicians from both Alzheimer centers.

The control participants were screened to exclude memory complaints, drug or alcohol abuse, major psychiatric disorders, and neurological or cerebrovascular diseases. They underwent an assessment including medical history, physical examination, extensive neuropsychological assessment, and an MRI of the brain, comparable to the work-up of patients.

All study participants underwent extensive neuropsychological assessment and MRI scanning at baseline and follow-up (Supplementary Table 1). Mean interval between the first visit (baseline measurement) and second visit (follow-up measurement) was 1.8 years (1.79 years for AD patients, 1.76 years for bvFTD patients, and 1.77 years for controls).
This study was performed in compliance with the Code of Ethics of the World Medical Association (Declaration of Helsinki). Ethical approval was obtained from the local ethics committees. Written informed consent from all participants was obtained.

Data acquisition

All participants underwent an MRI of the brain, on a 3 Tesla scanner using a standard 8-channel head coil, either in the VU University Medical Center (Signa HDxt, GE Healthcare, Milwaukee, WI, USA), or in the Leiden University Medical Center (Achieva, Philips Medical Systems, Best, The Netherlands) at baseline and follow-up.

Resting state fMRI T2*-weighted scans were acquired using whole brain multislice gradient echo planar imaging. Imaging parameters in the VU University Medical Center were: TR = 1.8 sec, TE = 35 msec, flip angle = 80°, 34 slices, resulting in a voxel size of 3.30 x 3.30 x 3.30 mm, including 10% interslice gap, 200 volumes, scan duration 6 minutes. Imaging parameters in the Leiden University Medical Center were: TR = 2.2 sec, TE = 30 msec, flip angle = 80°, 38 slices, resulting in a voxel size of 2.75 x 2.75 x 2.99 mm, including 10% interslice gap, 200 volumes, scan duration 7 minutes and 33 seconds. Participants were instructed to lie still with their eyes closed and not to fall asleep during the resting state scan.

Three-dimensional T1-weighted anatomical images were acquired, for registration purposes and to study group differences in gray matter volume. Imaging parameters in the VU University Medical Center were: TR = 7.8 msec, TE = 3 msec, flip angle = 12°, 180 slices, resulting in a voxel size of 0.98 x 0.98 x 1.00 mm. Imaging parameters in the Leiden University Medical Center were: TR = 9.8 msec, TE = 4.6 msec, flip angle = 8°, 140 slices, resulting in a voxel size of 0.88 x 0.88 x 1.20 mm. In the Leiden University Medical Center an additional high-resolution echo planar imaging scan was acquired for registration purposes (TR = 2.2 sec, TE = 30 msec, flip angle = 80°, 84 slices, resulting in a voxel size of 1.96 x 1.96 x 2.00 mm, including 10% interslice gap).
In addition, the MRI protocol included a 3D Fluid Attenuated Inversion Recovery (FLAIR) sequence, dual-echo T2-weighted sequence, and susceptibility weighted imaging (SWI) which were reviewed for brain pathology other than atrophy by an experienced radiologist. White matter hyperintensities were rated using the Fazekas scale [23]. Subjects with a score higher than 2 were excluded.

Data analysis

Before analysis, all MRI scans were submitted to a visual quality control check to ensure that no gross artifacts were present in the data. Data analysis was performed with Functional Magnetic Resonance Imaging of the Brain Software Library (FSL 5.0.1, Oxford, United Kingdom) [24]. Anatomical regions were determined using the Harvard-Oxford cortical and subcortical structures atlas integrated in FSL.

Gray matter volume

Structural MRI scans were analyzed with voxel-based morphometry (VBM) analysis [25] to study group differences and longitudinal changes in gray matter volume. First, the structural images were brain extracted and tissue-type segmented [26]. The resulting gray matter partial volume images were aligned to the gray matter MNI-152 standard space image (Montreal Neurological Institute, Montreal, QC, Canada) [27], followed by non-linear registration [28]. The images were averaged to create a study-specific template. Next, all native gray matter images were non-linearly registered to this study-specific gray matter template [25,29]. To correct for the contractions and enlargements due to the non-linear registration, each voxel of each registered gray matter image was multiplied by the Jacobian of the warp field, which defines the direction and the amount of modulation. The modulated segmented images were spatially smoothed with an isotropic Gaussian kernel with a full width at half maximum of 7 mm.

We studied 1) cross-sectional group differences at baseline (Fig. 1.1), 2) cross-sectional group differences at follow-up (Fig. 1.2), 3) longitudinal changes in gray matter volume within groups (i.e., within each group, the gray matter volume at follow-up different from that at baseline (delta))?
(Fig. 1.3). 4) group differences in longitudinal changes in gray matter volume (i.e., are the deltas different between groups?) (Fig. 1.4). To study group differences in gray matter volume at baseline and at follow-up, a general linear model (GLM) approach using analysis of variance F-tests with post hoc Bonferroni adjusted t-tests was applied. Age, gender, study center, and duration of symptoms were included as covariate in the statistical model. To study longitudinal changes in gray matter volume, individual gray matter maps from the second visit (follow-up) were subtracted from the corresponding gray matter maps from the first visit (baseline). This results for each subject in a map containing the differences in gray matter volume between the two time points (delta). These maps were concatenated across subjects into single 4D maps and were submitted to voxel-based statistical testing to study longitudinal changes in gray matter volume within (Fig. 1.3) and between groups (Fig. 1.4). F-tests were used adjusted for age, gender, study center, and duration of symptoms with post hoc t-tests. Voxel-wise non-parametric permutation testing [30] with 5000 permutations was performed using FSL-randomise correcting for multiple comparisons across voxels (statistical threshold set at p < 0.05, Family-Wise Error (FWE) corrected), using the Threshold-Free Cluster Enhancement (TFCE) technique [31].

Preprocessing of resting state fMRI data

The preprocessing of the resting state data consisted of motion correction [27], brain extraction [32], spatial smoothing using a Gaussian kernel with a full width at half maximum of 3 mm, and high-pass temporal filtering (cutoff frequency of 0.01 Hz). After preprocessing, the functional images were registered to the corresponding T1-weighted images using Boundary-Based Registration [33]. T1-weighted images were registered to the 2 mm isotropic MNI-152 standard space image (Montreal Neurological Institute, Montreal, QC, Canada) using nonlinear registration [34] with a warp resolution of 10 mm. High-resolution echo planar images (only available for subjects scanned in the Leiden University Medical Center) were used for an additional registration step between functional images and T1-weighted images. In order to achieve better comparison across voxels, subjects, time points, and centers, standardization on a voxel-by-voxel basis has been recommended [35]. We used the Z-standardization approach in which individual resting state fMRI time series were normalised
(standardized to z-scores) on a voxel-by-voxel basis using the mean and standard deviation of each individual resting state signal across time (previously described in [35]). Single-session independent component analysis (ICA) was performed on the preprocessed resting state data to decompose the data into distinct components for denoising purposes [36]. FMRIB’s ICA-based Xnoiseifier 1.05 (FIX) was used to auto-classify ICA components into “good” (i.e., functional signal) and “bad” (i.e., noise) components [37]. FIX removed unique variance related to “noise” components and motion confounds from the preprocessed fMRI data to denoise the resting state data and to increase the signal-to-noise ratio.

Resting state networks of interest

Group differences and longitudinal changes in functional connectivity were studied using the dual regression method of FSL (previously described in [38]). We used standard resting state networks as a reference to study functional connectivity in a standardized way [39,40]. Resting state functional connectivity was determined in terms of similarity of the BOLD fluctuations in the brain in relation to characteristic fluctuations in the predefined resting state networks [41,42]. These standardized resting state networks parcellate the brain into nine templates: network I) calcarine sulcus, precuneal cortex, and primary visual cortex (medial visual network); network II) superior and fusiform areas of lateral occipital cortex (lateral visual network); network III) superior temporal cortex, insular cortex, anterior cingulate cortex, auditory cortex, operculum, somatosensory cortices, thalamus (auditory system network); network IV) precentral and postcentral somatosensory somatomotor areas (sensorimotor system network); network V) rostral medial prefrontal cortex, precuneal cortex, posterior cingulate cortex (default mode network); network VI) medial and inferior prefrontal cortex, anterior cingulate and paracingulate gyri, prefrontal cortex (executive control network); networks VII and VIII) frontal pole, dorsolateral prefrontal cortex, parietal lobule, paracingulate gyrus, posterior cingulate cortex (right and left frontoparietal network), additionally we studied connectivity with network IX) anterior cingulate cortex, insular cortex, frontal orbital cortex, and frontal pole (salience network) (for further details see Fig. 3A and [42–44]). To account for noise, even after FIX, white matter and cerebrospinal fluid templates were included in the analysis [45–47].

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In the dual regression, individual time series were first extracted for each template, using the resting state networks [42,44] and the two additional white matter and cerebrospinal fluid maps [45–47], in a spatial regression against the individual fMRI data set (regression 1). The resulting matrices described temporal dynamics for each template and individual. Next, the temporal regressors were used to fit a linear model to the individual fMRI data set (regression 2), to estimate the spatial maps for each individual. This results in 3D images for each individual, with voxel-wise z-scores representing the functional connectivity to each of the templates. The higher the absolute value of the z-score, the stronger the connectivity to a network. Here, we studied 1) mean network connectivity (z-score) within the resting state networks of interest, and 2) network-to-region connectivity using a more detailed voxel-based analysis.

Mean network connectivity

First, we performed a whole-network analysis to study mean functional connectivity within each network of interest. For each participant, mean functional connectivity (z-score) per network was calculated. Figure 1 shows the statistical analyses that were performed. We studied 1) cross-sectional group differences at baseline (Fig. 1.1), 2) cross-sectional group differences at follow-up (Fig. 1.2), 3) longitudinal changes in connectivity within groups (i.e., is, within each group, the functional connectivity at follow-up different from that at baseline (delta)?) (Fig. 1.3), 4) group differences in longitudinal changes in connectivity (i.e., are the deltas different between groups?) (Fig. 1.4). We used analysis of covariance (ANCOVA; general linear model (GLM) procedure), adjusted for age, gender, study center, and duration of symptoms with post hoc Bonferroni adjusted t-tests (IBM SPSS Statistics Version 20, IBM Corp. Somers, NY, USA), to study cross-sectional group differences in mean network connectivity at baseline and at follow-up. We performed these analyses both without and with a correction for total gray matter volume of each network (at baseline or follow-up). To study longitudinal changes in mean network connectivity, individual mean z-score per network from the second visit (follow-up) were subtracted from the corresponding mean z-score from the first visit (baseline). This results in a delta score between the two time points per network for each subject. These delta scores were tested using an ANCOVA, adjusted for age, gender, study center, duration of
symptoms, and network connectivity at baseline, with post hoc Bonferroni adjusted t-tests, to study longitudinal changes in network connectivity within and between groups. These analyses were performed both without and with a correction for longitudinal changes in total gray matter volume of each network. Statistical threshold was set at $p < 0.05$ for all statistical tests.

Network-to-region connectivity

In a second, more detailed voxel-based analysis, we studied functional connectivity between resting state networks and localized brain regions. We used a GLM approach, as implemented in FSL, using F-tests, adjusted for age, gender, study center, and duration of symptoms with post hoc t-tests, to study cross-sectional group differences at baseline (Fig. 1.1) and at follow-up (Fig. 1.2). We performed these analyses both without and with a correction for voxel-based gray matter volume (at baseline or follow-up). To study longitudinal changes in network-to-region connectivity, individual functional connectivity maps (parameter estimates) from the second visit (follow-up) were subtracted from the corresponding functional connectivity maps from the first visit (baseline). This results for each subject in a map containing, per network, the differences in functional connectivity between the two time points (delta). These maps were concatenated across subjects into single 4D maps (one per predefined network) and were submitted to voxel-based statistical testing to study longitudinal changes in network-to-region connectivity within (Fig. 1.3) and between groups (Fig. 1.4). F-tests were used adjusted for age, gender, study center, and duration of symptoms with post hoc t-tests. These analyses were performed both without and with a correction for longitudinal voxel-based gray matter changes (i.e., delta gray matter volume). Per network, the statistical analyses were masked by the baseline one-sample map from the control group for that network. Voxel-wise non-parametric permutation testing [30] with 5000 permutations was performed using FSL-randomise correcting for multiple comparisons across voxels (statistical threshold set at $p < 0.05$, Family-Wise Error (FWE) corrected), using the Threshold-Free Cluster Enhancement (TFCE) technique [31].
Associations with cognitive performance

In a final analysis, we investigated the possible associations between changes in network-to-region connectivity and longitudinal changes in cognitive performance (i.e., Mini Mental State Examination (MMSE) score [48] and Frontal Assessment Battery (FAB) scores [49]), using linear regression analyses (IBM SPSS Statistics Version 20, IBM Corp. Somers, NY, USA), adjusted for age, gender, study center, and duration of symptoms (statistical threshold was set at p < 0.05).
**Results**

Demographic characteristics

Demographic data for all participants are summarized in Table 1. There were no differences between patient groups with regard to age at baseline, follow-up time, gender, study center distribution, level of education, and duration of symptoms. As expected, both dementia groups performed worse on cognitive tests compared with controls (all p < 0.05). Patients with AD performed worse at follow-up compared with baseline on MMSE (p = 0.016) and FAB (p = 0.049). Results of extensive neuropsychological assessment are shown in Supplementary Table 1.

Gray matter volume

Voxel-wise structural analysis at baseline revealed group differences in gray matter volume (Fig. 2A-B). Patients with AD showed less gray matter in posterior cingulate cortex, precuneal cortex, frontal medial cortex, temporal gyrus, hippocampus, and frontal orbital cortex compared with controls (Fig. 2A). Patients with bvFTD showed less gray matter in anterior cingulate cortex, insular cortex, and temporal pole compared with controls (Fig. 2B). No baseline differences in gray matter volume were found between patients with AD and patients with bvFTD. These differences in gray matter volume were also found at follow-up (Fig. 2C-D). **Longitudinal gray matter analyses showed over time decreased gray matter in the precuneal cortex, posterior cingulate cortex, temporal gyrus, parahippocampal gyrus, and accumbens in the AD group (Fig. 2E) and in the insular cortex, anterior cingulate cortex, and temporal gyrus in the bvFTD group (Fig. 2F).**

Mean network connectivity

We performed a whole-network analysis to study mean network functional connectivity in the nine resting state networks of interest (Fig. 3A). **These analyses were performed both without and with correction for total gray matter volume of each network.**
At baseline (Fig. 1.1), we found significant group differences in network VII and VIII. The results of post hoc testing showed lower mean connectivity in the right and left frontoparietal network (which include parietal lobule, paracingulate and postcingulate gyrus, and frontal pole) in the AD group compared with the control group (p = 0.045, p = 0.008, both uncorrected for gray matter volume) (Fig. 3B). When a correction for gray matter volume was applied functional connectivity in network VII (right frontoparietal network) was lower in the AD group compared with the control group (p = 0.024, gray matter corrected). No baseline differences in mean network connectivity were found in the bvFTD group.

At follow-up (Fig. 1.2), significant group differences were found in mean network connectivity in network VII. Post hoc testing showed lower connectivity in this right frontoparietal network in AD compared with controls (p = 0.010, uncorrected for gray matter volume) and in bvFTD compared with controls (p = 0.012, uncorrected for gray matter volume) (Fig. 3B). When a correction for gray matter volume was applied functional connectivity in network VII (right frontoparietal network) was lower in the AD group compared with the control group (p = 0.005, gray matter corrected) and in bvFTD compared with controls (p = 0.009, gray matter corrected).

When longitudinal changes in mean network connectivity were studied (Fig. 1.3), we found decreased mean connectivity in network VIII (left frontoparietal network) after the 1.8-year follow-up period in the bvFTD group (p = 0.021, uncorrected for gray matter volume) (Fig. 3B). No longitudinal changes in the bvFTD group were found when a gray matter volume correction was applied. In the AD and control group, no longitudinal changes in mean network connectivity were found.

When group differences in longitudinal changes were studied (Fig. 1.4), we found with post hoc tests group differences in longitudinal changes in mean network connectivity in network VII (right frontoparietal network) between AD and controls (p = 0.041, uncorrected for gray matter volume), and between bvFTD and controls (p = 0.043, uncorrected for gray matter volume). When a correction for gray matter volume was applied, longitudinal changes in mean network connectivity were found.
in network VII (right frontoparietal network) between AD and controls (p = 0.031, gray matter corrected), and between bvFTD and controls (p = 0.043, gray matter corrected). The mean connectivity in this network decreased more in AD and bvFTD patients than in controls.

Network-to-region connectivity

In a more detailed voxel-based analysis, we studied functional connectivity between the nine resting state networks of interest (Fig. 3A) and localized brain regions (Fig. 4).

At baseline (Fig. 1.1), we found significant group differences in network-to-region connectivity within four resting state networks: networks I, III, VI, and IX (Fig 4A, indicated in light orange). The results of post hoc testing showed higher functional connectivity in AD compared with controls between the lingual gyrus and network I (medial visual network), between the central opercular gyrus and network III (auditory system network), between the paracingulate gyrus and network VI (executive control network), and between the paracingulate gyrus and network IX (salience network) (Fig. 4A, AD > HC, yellow-to-red voxels, Table 2). No baseline differences in network-to-region connectivity were found in the bvFTD group. When a voxel-based correction for gray matter volume was applied comparable group differences in functional connectivity were found (Supplementary Figure 1, and Table 2 right column).

At follow-up (Fig. 1.2), significant group differences in network-to-region connectivity were found in six resting state networks: networks I, III, V, VI, VII, IX (Fig. 4B, indicated in light orange). Post hoc testing showed that the baseline differences in functional connectivity between AD patients and controls were also present at the 1.8-year follow-up measurement (Fig. 4B, AD > HC, yellow-to-red voxels). Note the lateralization of the effect in network III at follow-up. This lateralization was not visible in the images that were not corrected for FWE (uncorrected images not shown). The results of post hoc testing showed also group differences that were only present at follow-up: functional connectivity was lower in AD compared with controls between the angular gyrus and network VII (right frontoparietal network) (Fig. 4B, AD < HC, light-to-dark blue voxels), and in bvFTD between
the paracingulate gyrus and network V (default mode network), and between the angular gyrus and network VII (right frontoparietal network) (Fig. 4B, FTD < HC, light-to-dark blue voxels, Table 2). Functional connectivity in bvFTD was lower compared with AD patients between the lateral occipital cortex and network I (medial visual network), and between the anterior cingulate gyrus and network VI (executive control network) (Fig. 4B, FTD < AD, light-to-dark blue voxels, Table 2). After voxel-based correction for gray matter volume comparable group differences in functional connectivity were found (Supplementary Figure 1, and Table 2 right column).

When longitudinal changes in network-to-region connectivity were studied (Fig. 1.3), we found decreased connectivity over time within two networks: network VII and VIII (Fig. 5, indicated in light orange). In the AD group, functional connectivity between the precuneus and network VII (right frontoparietal network) decreased over time (Fig. 5A, and D). In the bvFTD group, functional connectivity between the supramarginal gyrus and network VII (right frontoparietal network), and between the inferior frontal gyrus and network VIII (left frontoparietal network) decreased over time (Fig. 5B-D). No longitudinal changes in network-to-region connectivity were found in the control group. After voxel-based correction for longitudinal gray matter changes comparable functional connectivity changes were found.

When group differences in longitudinal changes were studied (Fig. 1.4), we found group differences in longitudinal changes in network-to-region connectivity in network VII (Fig. 6, indicated in light orange). The results of post hoc testing showed group differences in longitudinal changes in connectivity between the supramarginal gyrus and network VII (right frontoparietal network) in AD (Fig. 6A, AD versus HC), and in bvFTD (Fig. 6B, FTD versus HC) when compared with controls. These small brain clusters show a decrease in functional connectivity over time in both patients groups, and an (insignificant) increase in the control group (Fig. 6C). Comparable functional connectivity changes were found when a voxel-based correction for longitudinal gray matter changes was applied.
Associations with cognitive performance

Finally, we studied whether the changes in network-to-region connectivity within groups (i.e. decreased connectivity with the precuneus in AD (Fig. 5A), and with the supramarginal (Fig. 5B) and inferior frontal gyrus (Fig. 5C) in bvFTD) were associated with changes in cognitive performance. In AD, no associations between changes in MMSE scores and changes in precuneal connectivity were found ($p = 0.894$). Changes in FAB scores were not associated with changes in supramarginal ($p = 0.310$) and inferior frontal gyrus ($p = 0.694$) connectivity in bvFTD.
Discussion

In this longitudinal study on resting state fMRI data in AD and bvFTD, we used whole-network and network-to-region analyses to study group differences in functional connectivity at baseline and at 1.8-year follow-up measurement, and changes in functional connectivity over time. We found disease-specific brain regions with longitudinal changes in functional connectivity in AD and bvFTD.

Over time, the connectivity between precuneus and right frontoparietal network decreased in AD, whereas in bvFTD connectivity between inferior frontal gyrus and left frontoparietal network decreased.

Our results suggest the potential of longitudinal resting state fMRI to delineate regions relevant for disease progression and for diagnostic accuracy, although the direct comparison between our relatively small AD and bvFTD groups did not yield significant group differences in longitudinal changes. Further studies, with larger patient groups, a longer follow-up time, and with more disease progression and neuropsychological decline, may give additional valuable information for disease progression and differential diagnosis.

This is the first study that investigates longitudinal changes in functional connectivity in both bvFTD and AD. In bvFTD, we found with the whole-network analysis decreasing longitudinal functional connectivity in the right and left frontoparietal network that encompassed frontal pole, dorsolateral prefrontal cortex, parietal lobule, paracingulate gyrus, and posterior cingulate cortex. In more detail, the network-to-region analysis showed decreased longitudinal connectivity between the left frontoparietal network and the inferior frontal gyrus. Cross-sectional differences in functional connectivity of the inferior frontal gyrus have been found between bvFTD and controls [21].

Furthermore, longitudinal connectivity between the supramarginal gyrus and the right frontoparietal network was decreased over time in bvFTD. The exact role of the supramarginal gyrus in the behavior of bvFTD is not clear. It has been shown that this brain area plays a crucial role in empathy [50], and
gray matter atrophy in the supramarginal gyrus has been reported in bvFTD [51], but is also common in AD [8]. In the present study, we found that in both bvFTD and AD the longitudinal changes in connectivity between the supramarginal gyrus and the right frontoparietal network were significantly different from those in control participants.

In AD, we found decreased longitudinal functional connectivity between the precuneus and the right frontoparietal network. The precuneus is particularly vulnerable for AD pathology, including gray matter atrophy and amyloid pathology [6,52]. Our finding is in line with the observation of decreasing longitudinal connectivity in the precuneus in AD [53].

In addition to the longitudinal changes, we reported cross-sectional group differences at two time points. Although connections with the average time series of the default mode network were not different between AD and controls, we did find group differences in specific regions of this network. Most prominent group differences between AD and controls were higher connectivity between cuneal cortex and the medial visual network, between the paracingulate gyrus and the executive control network, and between the paracingulate gyrus and the salience network. We were able to replicate our baseline findings 1.8 years later at the follow-up measurement. Visual inspection shows more extended group differences at the follow-up measurement, suggesting longitudinal changes in functional connectivity, however, these were not significant. Our baseline findings are in line with the observation of higher functional connectivity in the lingual gyrus, cuneal cortex, and salience network in AD [16,54,55], although not consistently observed [14,16]. While most studies reported lower functional connectivity in patients, there is evidence for higher functional connectivity as well [16,54,55].

In bvFTD, we observed lower functional connectivity compared with controls between the angular gyrus and the right frontoparietal network and between the paracingulate gyrus and the default mode network. These findings were comparable with a study that found differences in these functional connections when comparing patients with bvFTD and controls [21]. In our study, we found these
group differences only at follow-up, not at baseline. Furthermore, we were not able to find differences in the salience network connectivity in the bvFTD group. The inability to find baseline group differences in bvFTD is most likely related to the limited statistical power due to the relatively small number of subjects that was included in the bvFTD group. Another potential explanation is that we included less severely affected bvFTD patients, since we only included patients with scans available at both time points (baseline and follow-up). As a consequence, no data were available for the more severely affected subjects that dropped out of the study prematurely.

The most prominent finding in the direct cross-sectional comparison of AD and bvFTD is the lower functional connectivity between the anterior cingulate cortex and the executive control network in bvFTD compared with AD at the follow-up measurement. The anterior cingulate cortex is identified as a region that is among the first affected brain regions in bvFTD [56]. The deficits in social-emotional functions, which are common in bvFTD, rely on structures including the anterior cingulate cortex and frontoinsula [57,58]. Gray matter atrophy in these structures has shown to be more severe in bvFTD than in AD [59]. Cross-sectional group differences in the anterior cingulate cortex are observed as well in the two other resting state fMRI studies that performed the direct comparison of functional connectivity between patients with bvFTD and AD [16,20].

In addition to the longitudinal changes and group differences in functional connectivity, we found differences in gray matter volume between groups and changes in gray matter volume over time. When we correct the functional connectivity analyses for voxel-based gray matter volume, comparable group differences and longitudinal changes in functional connectivity were found, which might suggest that the functional connectivity differences are independent of gray matter volume.

The disease-specific changes in functional connectivity were not associated with changes in cognitive functioning. Lowest scores on a general measurement of cognitive performance (MMSE) [48] were found in patients with AD. As expected, we found over time a decline in MMSE scores in the AD group. Lowest scores on a general measurement of executive functioning (FAB) [49] were found in
the bvFTD group. Although we expect a decline in FAB scores in the bvFTD group, we found this decline in the AD group. AD patients showed low scores in the memory domain, but also in the executive functioning domain. Although executive functioning could be useful to differentiate AD from bvFTD [60,61], executive functioning, measured with FAB, may not discriminate both types of dementia [62,63]. It has been suggested that testing multiple cognitive domains is required to differentiate both types of dementia rather than focus on one cognitive test [63]. This might contribute to the lack of associations between one single measurements of cognitive functioning and changes in functional connectivity. In the current study, diagnoses were established according to the core clinical criteria for probable AD [2] and for bvFTD [3] and were therefore not based on one single neuropsychological test score.

The diagnosis FTD or AD can only be confirmed by post-mortem brain autopsy after death. A limitation of our study is that postmortem data were not available, therefore the possibility of misdiagnosis of the patients cannot be excluded. Nevertheless, all patients underwent an extensive dementia screening and were evaluated in a multidisciplinary panel including clinicians from different centers specialized in dementia. Only dementia patients that fulfilled the most recent clinical criteria for probable AD [2] and bvFTD [3] were included in the present study. Another limitation of our study is the relatively small number of subjects that was included in the bvFTD group. Further studies, with larger patient groups and more disease progression and neuropsychological decline, may give additional valuable information. A further limitation of the current study includes the possible effects of medication on the functional connections in the AD and bvFTD groups.

This longitudinal study showed data that were collected in two centers. The main strength of multicenter studies is the increased generalizability of the study findings. However, multicenter studies have also its limitations, since the data will be less homogeneous than in single center studies. To increase homogeneity between centers in the current study, we evaluated all patients in a multidisciplinary panel including clinicians from different centers specialized in dementia, we used a standardization approach in order to achieve better comparison across voxels, subjects, and centers.
[35], and we added center as covariate in all statistical models, following previous approaches [16,64].

To conclude, we used longitudinal resting state fMRI data of patients with AD and patients with bvFTD and found disease-specific brain regions with longitudinal connectivity changes. Over time, **functional connectivity between precuneus and the right frontoparietal network** decreased in AD, and **connectivity between the inferior frontal gyrus and left frontoparietal network** decreased in bvFTD. This suggests the potential of longitudinal resting state fMRI to delineate regions relevant for disease progression and for diagnostic accuracy, although we did not find group differences in longitudinal changes in the direct comparison of AD and bvFTD patients.
Acknowledgements

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H. Vrenken has received research support from Merck-Serono, Novartis, and Pfizer, and speaker honoraria from Novartis. All funds were paid to his institution.

F. Barkhof serves / has served on the advisory boards of: Bayer-Schering Pharma, Sanofi-Aventis, Biogen Idec, UCB, Merck-Serono, Novartis, and Roche. He has been a speaker at symposia organized by the Serono Symposia Foundation. For all his activities he receives no personal compensation.

P. Scheltens serves / has served on the advisory boards of: Genentech, Novartis, Roche, Danone, Nutricia, Baxter and Lundbeck. He has been a speaker at symposia organized by Lundbeck, Merz, Danone, Novartis, Roche and Genentech. He serves on the editorial board of Alzheimer’s Research & Therapy and Alzheimer’s Disease and Associated Disorders. And he is a member of the scientific advisory board of the EU Joint Programming Initiative and the French National Plan Alzheimer. For all his activities he receives no personal compensation.
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Table 1  Characteristics of the study population

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>AD (n=20)</th>
<th>bvFTD (n=12)</th>
<th>HC (n=22)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>65.1 (8.0)</td>
<td>64.7 (5.4)</td>
<td>63.8 (5.0)</td>
</tr>
<tr>
<td>Follow-up time (years)</td>
<td>1.79 (0.43)</td>
<td>1.76 (0.43)</td>
<td>1.77 (0.59)</td>
</tr>
<tr>
<td>Gender (male/female)</td>
<td>16/4</td>
<td>9/3</td>
<td>14/8</td>
</tr>
<tr>
<td>Study center (VUMC/LUMC)</td>
<td>16/4</td>
<td>8/4</td>
<td>12/10</td>
</tr>
<tr>
<td>Level of educationb</td>
<td>4.9 (1.1)</td>
<td>5.0 (1.3)</td>
<td>5.6 (0.7)</td>
</tr>
<tr>
<td>Duration of symptoms (median ± min/max) (months)</td>
<td>36 ± 12/108</td>
<td>60 ± 12/192</td>
<td>n/a</td>
</tr>
<tr>
<td>MMSE (max score: 30)</td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Baseline</td>
</tr>
<tr>
<td></td>
<td>23.2 (2.9)</td>
<td>19.8 (5.2)\textsuperscript{i}</td>
<td>24.3 (3.9)</td>
</tr>
<tr>
<td>FAB (max score: 18)</td>
<td>14.1 (2.7)</td>
<td>11.7 (3.7)\textsuperscript{i}</td>
<td>13.3 (2.7)</td>
</tr>
<tr>
<td>CDR (max score: 3)</td>
<td>1.1 (0.8)</td>
<td>1.2 (0.5)</td>
<td>1.3 (0.8)</td>
</tr>
<tr>
<td>GDS (max score: 15)</td>
<td>2.7 (3.4)</td>
<td>1.9 (2.0)</td>
<td>1.4 (0.8)</td>
</tr>
</tbody>
</table>

Abbreviations: AD = Alzheimer’s disease; bvFTD = behavioral variant of frontotemporal dementia; HC = healthy controls; MMSE = Mini-Mental State Examination; FAB = Frontal Assessment Battery; CDR = Clinical Dementia Rating Scale; GDS = Geriatric Depression Scale

Values are means (standard deviation) for continuous variables or numbers for dichotomous variables. Scores on FAB and GDS were missing in 5 patients.

\textsuperscript{i}Imaging was performed either in the Alzheimer Center of the VU University Medical center (VUMC) or in the Leiden University Medical Center (LUMC) in the Netherlands.

\textsuperscript{b}Level of education was determined on a Dutch 7-point scale ranging from 1 (less than elementary school) to 7 (university or technical college).

\textsuperscript{i}Significant differences between baseline and follow-up
Table 2  Group differences in network-to-region connectivity at baseline and at follow-up

<table>
<thead>
<tr>
<th>Network and contrast</th>
<th>Brain structurea</th>
<th>Side</th>
<th>Peak coordinates (MNI)</th>
<th>Peak T score</th>
<th>GM uncorrected</th>
<th>GM corrected</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>x    y    z</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Group differences at baseline</strong></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Medial visual network (I)</td>
<td>Lingual gyrus</td>
<td>R</td>
<td>10    -74   0</td>
<td>4.32</td>
<td>5.49</td>
<td></td>
</tr>
<tr>
<td>AD &gt; HC</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Auditory system network (III)</td>
<td>Central opercular cortex</td>
<td>R</td>
<td>56    4    2</td>
<td>4.51</td>
<td>5.16</td>
<td></td>
</tr>
<tr>
<td>AD &gt; HC</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Executive control network (VI)</td>
<td>Paracingulate gyrus</td>
<td>L</td>
<td>-12   44    18</td>
<td>4.23</td>
<td>4.73</td>
<td></td>
</tr>
<tr>
<td>AD &gt; HC</td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Salience network (IX)</td>
<td>Paracingulate gyrus</td>
<td>L</td>
<td>10    20    30</td>
<td>4.26</td>
<td>3.50</td>
<td></td>
</tr>
<tr>
<td>AD &gt; HC</td>
<td></td>
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<td></td>
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<td></td>
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</tr>
<tr>
<td><strong>Group differences at follow-up</strong></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Medial visual network (I)</td>
<td>Lingual gyrus</td>
<td>R</td>
<td>18    -70   -4</td>
<td>4.82</td>
<td>5.34</td>
<td></td>
</tr>
<tr>
<td>AD &gt; HC</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Auditory system network (III)</td>
<td>Cuneal cortex</td>
<td>R</td>
<td>18    -80   28</td>
<td>3.87</td>
<td>3.87</td>
<td></td>
</tr>
<tr>
<td>AD &gt; HC</td>
<td></td>
<td></td>
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<tr>
<td>Executive control network (VI)</td>
<td>Planum polare</td>
<td>L</td>
<td>-48   -14   -4</td>
<td>5.24</td>
<td>4.90</td>
<td></td>
</tr>
<tr>
<td>AD &gt; HC</td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Right frontoparietal network (VII)</td>
<td>Angular gyrus</td>
<td>R</td>
<td>58    -52   50</td>
<td>4.54</td>
<td>3.63</td>
<td></td>
</tr>
<tr>
<td>AD &lt; HC</td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Salience network (IX)</td>
<td>Paracingulate gyrus</td>
<td>R</td>
<td>4     42    30</td>
<td>5.32</td>
<td>5.32</td>
<td></td>
</tr>
<tr>
<td>AD &gt; HC</td>
<td>Anterior cingulate gyrus</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Default mode network (V)</td>
<td>Paracingulate gyrus</td>
<td>R</td>
<td>10    52    -4</td>
<td>4.48</td>
<td>3.87</td>
<td></td>
</tr>
<tr>
<td>bvFTD &lt; HC</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Right frontoparietal network (VII)</td>
<td>Angular gyrus</td>
<td>R</td>
<td>56    -52   48</td>
<td>4.47</td>
<td>3.76</td>
<td></td>
</tr>
<tr>
<td>bvFTD &lt; HC</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Medial visual network (I)</td>
<td>Lateral occipital cortex</td>
<td>R</td>
<td>24    -72   28</td>
<td>4.50</td>
<td>3.80</td>
<td></td>
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<tr>
<td>bvFTD &lt; AD</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Executive control network (VI)</td>
<td>Anterior cingulate gyrus</td>
<td>R</td>
<td>6     -10   38</td>
<td>4.61</td>
<td>4.41</td>
<td></td>
</tr>
<tr>
<td>bvFTD &lt; AD</td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>
Abbreviations: MNI = Montreal Neurological Institute Montreal, QC, Canada; GM = gray matter; AD = Alzheimer’s disease; HC = healthy controls; bvFTD = behavioral variant frontotemporal dementia; R = right; L = left

*Full list of structures with group differences in network-to-region connectivity (Fig. 4). Between group effects are independent of physiological noise, age, gender, study center, duration of symptoms. Analyses were performed with and without voxel-based gray matter (GM) volume correction (two right columns). Thresholding using p < 0.05, FWE corrected, based on the TFCE statistic image. For each peak voxel x-, y-, and z-coordinates in the MNI-152 standard space image are given.*
## Supplementary Table 1  Neuropsychological performance

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>AD (n=20)</th>
<th>bvFTD (n=12)</th>
<th>HC (n=22)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>MMSE</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>max score 30</td>
<td>23.2 (2.9)</td>
<td>24.3 (3.9)</td>
<td>29.1 (0.9)</td>
</tr>
<tr>
<td><strong>FAB</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>max score 18</td>
<td>14.1 (2.7)</td>
<td>13.3 (2.7)</td>
<td>17.4 (1.2)</td>
</tr>
<tr>
<td><strong>CDR</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>max score 3</td>
<td>1.1 (0.8)</td>
<td>1.3 (0.8)</td>
<td>0.0 (0.0)</td>
</tr>
<tr>
<td><strong>GDS</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>max score 15</td>
<td>1.2 (0.5)</td>
<td>1.7 (0.8)</td>
<td>0.0 (0.0)</td>
</tr>
<tr>
<td><strong>MMSE</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>max score 75</td>
<td>24.3 (3.9)</td>
<td>29.1 (0.9)</td>
<td>29.0 (1.5)</td>
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<tr>
<td><strong>FAB</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>max score 18</td>
<td>10.5 (6.3)</td>
<td>17.2 (1.1)</td>
<td></td>
</tr>
<tr>
<td><strong>CDR</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>max score 3</td>
<td>0.0 (0.0)</td>
<td>0.0 (0.0)</td>
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</tr>
<tr>
<td><strong>GDS</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>max score 15</td>
<td>0.8 (1.4)</td>
<td>1.2 (1.2)</td>
<td></td>
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<tr>
<td><strong>RAVLT immediate recall</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>max score 75</td>
<td>49.0 (8.9)</td>
<td>45.4 (9.4)</td>
<td></td>
</tr>
<tr>
<td><strong>RAVLT delayed recall</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>max score 15</td>
<td>9.5 (2.9)</td>
<td>8.6 (2.8)</td>
<td></td>
</tr>
<tr>
<td><strong>VAT</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>max score 6</td>
<td>5.8 (0.4)</td>
<td>5.7 (0.4)</td>
<td></td>
</tr>
<tr>
<td><strong>Digit span, forward</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>max score 30</td>
<td>10.4 (4.0)</td>
<td>12.1 (3.6)</td>
<td></td>
</tr>
<tr>
<td><strong>Digit span, backward</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>max score 30</td>
<td>9.5 (2.9)</td>
<td>8.6 (2.8)</td>
<td></td>
</tr>
<tr>
<td><strong>TMT A</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>time in seconds</td>
<td>36.8 (14.8)</td>
<td>37.9 (15.4)</td>
<td></td>
</tr>
<tr>
<td><strong>TMT B</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>time in seconds</td>
<td>82.7 (31.8)</td>
<td>81.1 (29.2)</td>
<td></td>
</tr>
<tr>
<td><strong>Stroop I</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>time in seconds</td>
<td>46.7 (7.1)</td>
<td>47.5 (7.5)</td>
<td></td>
</tr>
<tr>
<td><strong>Stroop II</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>time in seconds</td>
<td>60.7 (9.7)</td>
<td>60.0 (8.7)</td>
<td></td>
</tr>
<tr>
<td><strong>Stroop III</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>time in seconds</td>
<td>97.3 (19.8)</td>
<td>94.9 (20.6)</td>
<td></td>
</tr>
<tr>
<td><strong>Categorical fluency</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>responses in 1 minute</td>
<td>22.6 (5.4)</td>
<td>23.2 (6.6)</td>
<td></td>
</tr>
<tr>
<td><strong>Letter fluency</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>responses in 1 minute</td>
<td>12.6 (4.6)</td>
<td>14.3 (3.8)</td>
<td></td>
</tr>
<tr>
<td><strong>LDST</strong></td>
<td>Baseline</td>
<td>Follow-up</td>
<td>Follow-up</td>
</tr>
<tr>
<td>responses in 1 minute</td>
<td>34.1 (6.5)</td>
<td>34.7 (6.0)</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: AD = Alzheimer’s disease; bvFTD = behavioral variant frontotemporal dementia; HC = healthy controls; MMSE = Mini-Mental State Examination; FAB = Frontal Assessment Battery; CDR = Clinical Dementia Rating Scale; GDS = Geriatric Depression Scale; RAVLT = Rey Auditory Verbal Learning Test; VAT = Visual Association Test; TMT = Trail Making Test; LDST = Letter Digit Substitution Test
Significant differences between baseline and follow-up

Significant differences between AD and bvFTD
**Figure legends**

**Figure 1**  **The four main analyses of this study**

This hypothetical model is intended to show the four main analyses performed in the current study. Data points reflect examples of functional connectivity in patients with Alzheimer’s disease (AD), patients with behavioral variant frontotemporal dementia (FTD), and healthy controls (HC) at two time points (baseline and follow-up). We have studied 1) cross-sectional group differences at baseline, 2) cross-sectional group differences at follow-up, 3) longitudinal changes within groups (i.e., is functional connectivity at follow-up different from that at baseline (delta)?), 4) group differences in longitudinal changes (i.e., are the deltas different between groups?).

**Figure 2**  **Group differences and longitudinal changes in gray matter volume**

Differences in gray matter volume between Alzheimer’s disease (AD), behavioral variant frontotemporal dementia (FTD), and healthy controls (HC) (TFCE, FWE-corrected, p < 0.01). Images are overlaid on the most informative sagittal, coronal, and axial slices of the MNI-152 standard anatomical image (x, y, and z coordinates of each slice are given). At baseline, A) patients with AD showed less gray matter in precuneal cortex, posterior cingulate cortex, frontal medial cortex, temporal gyrus, hippocampus, and frontal orbital cortex compared with controls and B) patients with bvFTD showed less gray matter in anterior cingulate cortex, insular cortex, and temporal pole compared with controls. These differences in gray matter volume were also found at follow-up (C-D).

*Longitudinal changes in gray matter volume were found in E) the precuneal cortex, posterior cingulate cortex, temporal gyrus, parahippocampal gyrus, and accumbens in the AD group and in F) the insular cortex, anterior cingulate cortex, and temporal gyrus in the bvFTD group.*

**Figure 3**  **Mean network connectivity in resting state networks of interest**

A) Spatial maps of eight predefined resting state networks of interest. Images are overlaid on the most informative sagittal, coronal, and axial slices of the MNI-152 standard anatomical image (x, y, and z coordinates of each slice are given). Images are displayed following the radiological convention,
which means that the left side of the image corresponds to the right hemisphere and vice versa. B) Bar graphs show mean ($\pm$ standard error) functional connectivity within each resting state network for patients with Alzheimer’s disease (AD, blue), patients with behavioral variant frontotemporal dementia (FTD, red), and healthy controls (HC, green) both at baseline and follow-up. Asterisks indicate significant group differences or longitudinal changes (post hoc t-tests, Bonferroni corrected).

**Figure 4**

**Group differences in network-to-region connectivity – uncorrected for gray matter volume**

Cross-sectional group differences in functional connectivity between resting state networks (indicated in light orange) and localized brain regions (indicated in yellow-to-red and light-to-dark blue, full list of structures in Table 2), **uncorrected for voxel-based gray matter volume** (see Supplementary Figure 1 for results corrected for voxel-based gray matter volume). A) Post hoc testing showed at baseline higher connectivity in Alzheimer’s disease compared with controls (AD > HC, indicated in yellow-to-red). B) The results of post hoc testing showed that the baseline differences were also present at 1.8-year follow-up measurement. At follow-up, functional connectivity was lower in behavioral variant frontotemporal dementia compared with controls (FTD < HC, indicated in light-to-dark blue), and compared with AD patients (FTD < AD, indicated in light-to-dark blue). F-tests did not show at baseline group differences in networks I, V, VI, VII, which is for illustration purposes illustrated by brains without yellow-to-red and light-to-dark blue colored voxels. P values are color coded from 0.05 FWE corrected (red / dark-blue) to < 0.0001 FWE corrected (yellow / light-blue), **uncorrected for gray matter volume**. Images are overlaid on the sagittal, coronal, and transverse slices of the MNI-152 standard anatomical image (x, y, and z coordinates of each slice are given). Images are displayed following the radiological convention, which means that the left side of the image corresponds to the right hemisphere and vice versa.

**Figure 5**  

**Longitudinal changes in network-to-region connectivity within a group**

Longitudinal changes in functional connectivity between **the right frontoparietal network** (indicated in light orange) and A) precuneus in Alzheimer’s disease (AD, blue) and B) supramarginal gyrus (FTD,
red) and C) between the left frontoparietal network (indicated in light orange) inferior frontal gyrus in behavioral variant frontotemporal dementia (FTD, red) (uncorrected for gray matter volume). Images are overlaid on the MNI-152 standard anatomical image (coordinates of each slice are given). Images are displayed following the radiological convention, which means that the left side of the image corresponds to the right hemisphere and vice versa. D) Graphs are included to show the directionality of the effects. This shows decreased longitudinal functional connectivity in AD (blue) and FTD (red).

**Figure 6**  
**Group differences in longitudinal changes in network-to-region connectivity**

Group differences in longitudinal changes in functional connectivity between the right frontoparietal network (indicated in light orange and by roman numerals) and A) the supramarginal gyrus in patients with Alzheimer’s disease (AD, blue), and B) the supramarginal gyrus in patients with behavioral variant frontotemporal dementia (FTD, red) compared with healthy controls (HC) (uncorrected for gray matter volume). Images are overlaid on the MNI-152 standard anatomical image (coordinates of each slice are given). Images are displayed following the radiological convention, which means that the left side of the image corresponds to the right hemisphere and vice versa. C) Graphs are included to show the directionality of the effects. This shows group differences in changes in functional connectivity in patients with AD (blue) and in patients with FTD (red) compared with healthy controls (HC, green). The increase in functional connectivity over time in the healthy control group was not significant.

**Supplementary Figure 1**  
**Group differences in network-to-region connectivity – corrected for gray matter volume**

Cross-sectional group differences in functional connectivity between resting state networks (indicated in light orange) and localized brain regions (indicated in yellow-to-red and light-to-dark blue, full list of structures in Table 2), corrected for voxel-based gray matter (GM) volume. A) Post hoc testing showed at baseline higher connectivity in Alzheimer’s disease compared with controls (AD > HC, indicated in yellow-to-red). B) The results of post hoc testing showed that the baseline differences were also present at 1.8-year follow-up measurement. At follow-up, functional connectivity was
lower in behavioral variant frontotemporal dementia compared with controls (FTD < HC, indicated in light-to-dark blue), and compared with AD patients (FTD < AD, indicated in light-to-dark blue). P values are color coded from 0.05 FWE corrected (red / dark-blue) to < 0.0001 FWE corrected (yellow / light-blue), corrected for gray matter volume. Images are overlaid on the sagittal, coronal, and transverse slices of the MNI-152 standard anatomical image (x, y, and z coordinates of each slice are given).
Figure 1  The four main analyses of this study
Figure 2  Group differences and longitudinal changes in gray matter volume

A  Baseline group difference  AD < HC

B  Baseline group difference  FTD < HC

C  Follow-up group difference  AD < HC

D  Follow-up group difference  FTD < HC

E  Longitudinal change within AD group

F  Longitudinal change within FTD group
Figure 3: Mean network connectivity in resting state networks of interest

A

I
Medial visual network

II
Lateral visual network

III
Auditory system network

IV
Sensorimotor system network

V
Default mode network

VI
Executive control network

VII
Right frontoparietal network

VIII
Left frontoparietal network

IX
Salience network

B

Functional connectivity within resting state network (Z-score)

Baseline Follow-up

[A: Brain images showing different networks; B: Bar charts comparing functional connectivity across conditions]
Group differences in network-to-region connectivity - uncorrected for gray matter volume

Figure 4
Figure 5  Longitudinal changes in network-to-region connectivity within a group
Figure 6  Group differences in longitudinal changes in network-to-region connectivity

A  Right frontoparietal network (VII)  AD vs HC

B  Right frontoparietal network (VII)  FTD vs HC

C  Functional connectivity (z-score)

Baseline  Follow-up
Supplementary Figure 1

Group differences in network-to-region connectivity - corrected for gray matter volume