Functional Connectivity Alterationsof the Temporal Lobe and Hippocampus

- in Semantic Dementia and Alzheimer's
- Disease
- ⁵ Simon Schwab^{a,d}, Soroosh Afyouni^a, Yan Chen^b, Zaizhu Han^c, Qihao Guo^c, Thomas Dierks^d,
- Lars-Olof Wahlund^e and Matthias Grieder^{d,*}
- ^aBig Data Institute, Li Ka Shing Centre for Health Information and Discovery, Nuffield Department of Population
- Health, University of Oxford, Oxford, United Kingdom
- ^bState Key Laboratory of Cognitive Neuroscience and Learning, Beijing Normal University, Beijing, China
- ^cDepartment of Neurology, Huashan Hospital, Fudan University, Shanghai, China
- ¹¹ ^dTranslational Research Center, University Hospital of Psychiatry and Psychotherapy, University of Bern, Bern,
- 12 Switzerland

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- ¹³ ^eDepartment NVS, Karolinska Institute, Division of Clinical Geriatrics, Stockholm, Sweden
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15 Abstract.

- 16 Background: Semantic memory impairments in semantic dementia are attributed to atrophy and functional disruption of
- the anterior temporal lobes. In contrast, the posterior medial temporal neurodegeneration found in Alzheimer's disease
- is associated with episodic memory disturbance. The two dementia subtypes share hippocampal deterioration, despite a
- ¹⁹ relatively spared episodic memory in semantic dementia.
- **Objective:** To unravel mutual and divergent functional alterations in Alzheimer's disease and semantic dementia, we assessed functional dementiation of the dimensional disease (n 10) semantic dementiation of the alternative (n 22) and head the second disease (n 10) semantic dementiation of the second disease (n 10) semantic d
- functional connectivity between temporal lobe regions in Alzheimer's disease (n = 16), semantic dementia (n = 23), and healthy controls (n = 17).
- Methods: In an exploratory study, we used a functional parcellation of the temporal cortex to extract time series from 66
 regions for correlation analysis.
- Results: Apart from differing connections between Alzheimer's disease and semantic dementia that yielded reduced functional
- connectivity, we identified a common pathway between the right anterior temporal lobe and the right orbitofrontal cortex
- in both dementia subtypes. This disconnectivity might be related to social knowledge deficits as part of semantic memory
- 28 decline. However, such interpretations are preferably made in a holistic context of disease-specific semantic impairments and
- ²⁹ functional connectivity changes.

^{*}Correspondence to: Matthias Grieder, PhD, Translational Research Center, University Hospital of Psychiatry and Psychotherapy, Bolligenstrasse 111, 3000 Bern 60, Switzerland. Tel.: +41 319328351; Fax: +41 319309961; E-mail: matthias.grieder@upd.unibe.ch.

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34 Keywords: Alzheimer's disease, functional connectivity, semantic dementia, temporal lobe

30 INTRODUCTION

Everybody occasionally experiences difficulties in 31 integrating past events into an accurate context-a 32 condition classified as an episodic memory dis-33 turbance. Intact episodic memory [1] requires the 34 processing of information about chronology, place, 35 and the protagonists who were involved in an event. 36 The capability to store and retrieve autobiographi-37 cal memory is, however, not sufficient for an intact 38 episodic memory. Humans also strongly rely on 39 a fully functioning semantic memory. Concretely, 40 semantic memory reflects our general knowledge 41 about concepts such as objects, people, and words. 42 Thus, only a sound interplay of these two memory 43 systems, episodic and semantic memory, allows a 44 cognitively healthy state of an individual. 45

Previously two initially contradicting models of 46 the neurophysiological organization of semantic 47 memory have been harmonized as what can be char-48 acterized as a 'cortically distributed plus semantic 49 hub' theory [2, 3]. The term "distributed" refers to 50 the idea that regions which process semantic con-51 cepts receive multimodal input from corresponding 52 brain regions (e.g., visual attributes from visual brain 53 regions, tactile attributes from the sensorimotor cor-54 tex, etc.). Subsequently, these multimodal inputs 55 from distributed cortical areas converge to so-called 56 unitary semantic concepts in the semantic hub [4, 5]. 57 The semantic hub was found to be localized bilat-58 erally in the anterior temporal lobe, a region which 59 is atrophied and hypometabolized in patients with 60 the semantic variant of primary progressive aphasia, 61 also known as the temporal variant of frontotemporal 62 dementia (FTD) or semantic dementia (SD) [5-7]. In 63 SD, the onset of gray matter atrophy occurs in the 64 anterior temporal lobes, frequently with an asymme-65 try toward the more affected left hemisphere. With 66 progression of the disease, the temporal pole and 67 medial as well as lateral temporal areas are degen-68 erated [8]. However, the patients seem to exhibit an 69 almost intact episodic memory, when tested non-70 verbally, while their semantic memory is severely 71 deteriorated [9, 10]. 72

In contrast to SD, patients with Alzheimer's disease (AD) show predominantly episodic memory impairments, and semantic memory deficits can only be observed to a minor degree [11–13]. AD has been described as a disconnection syndrome, that is, connections of functionally or structurally linked brain regions that are part of a network become increasingly disrupted [14–16]. This degenerative mechanism has been associated with the cognitive deficits of patients with AD [17–19]. A common finding in AD is that gray matter atrophy onset can be localized in the hippocampal, posterior cingulate, and lateral parietal brain regions, as well as in the amygdala [20, 21]. The hippocampus forms a core region for episodic memory encoding. However, it has also been associated with semantic memory functions [22]. In fact, Burianova and colleagues [22] postulated that the hippocampus is part of a common declarative memory network, suggesting that the hippocampus has a key role in both semantic as well as episodic and autobiographical memory.

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The properties of functional systems, as for example Burianova and colleagues' proposed declarative memory network, are commonly assessed by the use of a resting-state functional connectivity (FC) analysis. The human resting-state is characterized by spatially discriminate brain regions that co-activate and deactivate at a low temporal frequency, commonly known as resting-state networks [23, 24]. These functional systems, or resting-state networks, are commonly assessed using blood-oxygen level dependent resting-state fMRI. It has become very popular to study FC alterations in various mental and neurological disorders including AD, demonstrating a relationship between disease and abnormalities in resting-state networks [25–27].

FC changes (i.e., decreases and increases of connectivity strengths) in AD have been found predominantly in the hippocampus and the default mode network [28–31]. With the progression of the disease, structural and functional connectivity distortions affect several networks, particularly those involving the para hippocampus [17, 32]. In SD, FC appears to be deteriorated in regions either affected

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by or proximate to the core of atrophy, located in
regions such as the temporal pole, anterior middle
temporal gyrus, inferior temporal gyrus, and insula
[6, 33–35]. Furthermore, reduced FC of the anterior
temporal lobe with various cortical regions was also
found in SD [2].

Considering these findings as well as the distinct 123 pathology of AD and SD, it is likely that the neuronal 124 loss of hippocampal cells that results in gray matter 125 atrophy certainly affects the functional networks in 126 a way that generates episodic memory deficits. Tem-127 poral pole atrophy alone might not be necessary (but 128 sufficient) to lead to semantic impairment. Follow-129 ing these findings, La Joie et al. [36] identified the 130 hippocampus as the 'main crossroad' between brain 131 networks that are disrupted in AD and SD. Despite the 132 growing body of research, the common and divergent 133 changes of FC among regions of the temporal lobes 134 in AD and SD are not fully understood. A caveat 135 when interpreting the existing literature is the com-136 mon use of anatomical/structural parcellation instead 137 of a functional parcellation to study FC. Functional 138 parcellations have the advantage that the resulting 139 functional regions of interest (ROIs) are homoge-140 neous, i.e., the voxels have similar time courses. On 141 the other hand, parcellations based on brain structure 142 can merge the time series across functionally different 143 areas which can be problematic [37]. 144

This proof-of-concept study aimed at disentan-145 gling FC alterations of the temporal lobe in AD and 146 SD using a refined division of temporal subregions: 147 sixty-six functional regions of interest (ROIs) of the 148 temporal lobes from a functional atlas [38]. In con-149 trast to numerous previous studies, we accounted for 150 structural changes (i.e., gray matter density) in order 151 to extract FC time series data from preserved gray 152 matter tissue which can still be functional [39, 40]. 153 In other words, results from the FC analysis reflect 154 the functional reorganization of the temporal lobes 155 affected by atrophy. 156

A common issue with studies involving patients 157 with SD is the small sample size due to the low 158 prevalence and relatively difficult diagnosis. In order 159 to overcome this to some extent, we pooled two 160 data sets from two different recording sites (see 161 Method section for details). Orban et al. [41] showed 162 the advantage of multisite fMRI-data in multivariate 163 fMRI analysis. Their approach appears to be gener-164 alizable; however, in our study, we were not able to 165 accomplish an evenly matched number of patients or 166 controls at each MRI scanner site, which is a prereq-167 uisite for a correct experimental design. In particular, 168

the circumstance that the majority of SD patients was scanned at the Shanghai site and all AD patients and healthy controls (HC) were scanned at the Stockholm site, increases the likelihood of false positive contrasts between the groups due to instrumental artifacts. Other inherent limitations will be addressed in the discussion section (e.g., site-specific diagnostic criteria, neuropsychological testing, and fMRI acquisition procedures).

Despite the exploratory analysis approach to test all possible connections, based on previous findings described above, the following hypotheses were tested: in AD, we expected FC alterations in the hippocampus, parahippocampal ROIs, and possibly posterior temporal ROIs. In SD, altered FC was anticipated in the hippocampus, the fusiform gyrus, and the temporal pole.

METHODS

Participants

We analyzed resting-state fMRI data from a total of 62 participants from three groups: semantic dementia (SD), Alzheimer's disease (AD), and a healthy elderly control group (HC). We examined all the functional MRI data and excluded six datasets due to insufficient data quality (see data quality control). The final sample consisted of 56 participants: Twenty-three patients with SD, with a mean age (\pm standard deviation) of 62 \pm 7.6, 16 patients with AD, mean age of 70 ± 8.5 , and 17 individuals in the HC group, mean age 70 ± 3.4 ; see Table 1 for demographics and clinical variables. Patients with SD from the Stockholm site (n=7) were recruited throughout Sweden and diagnosed using the criteria of Neary et al. [42], while patients with SD from Shanghai were recruited from Huashan Hospital in Shanghai (n=19), according to the criteria of Gorno-Tempini et al. [43]. The main diagnostic criteria of both guidelines share clinical observation features such as impaired word naming and comprehension, spared repetition, and surface dyslexia and dysgraphia. Differences in these two diagnostic criteria, as for instance the introduction of brain imaging as a supportive diagnostic feature in Gorno-Tempini et al. (2011), were not relevant, because also the Swedish patients underwent MRI to assess anterior temporal lobe atrophy. Patients with AD were recruited at the Memory Clinic of the Geriatric Department at Karolinska University Hospital in Huddinge, Sweden (n = 19). Their diagnosis was

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	HC $(n = 17)$	Normative data [†]	AD $(n = 16)$	SD (n=23)	
	Mean (std. dev.)	Mean (std. dev.)	Mean (std. dev.)	Mean (std. dev.)	р
Age, y	67.9 (3.3)		68.4 (8.5)	61.5 (7.4)	0.004
Gender (F:M)	12:5		7:9	10:13	-
Education, y	13.9 (3.1)		13.1 (3.0)	$12.4 (1.5)^3$	0.61
CDS	-		1.0 (1.0)	$1.8(2.2)^3$	0.60
GDS	-		2.9 (0.8)	$3.8(0.4)^3$	0.031
MMSE (max 30)	28.8 (0.8)		24.5 (4.8)	$20.8(5.2)^4$	< 0.0001
BNT (max 60)	54.4 (3.7)	54.0 (4.5)	45.6 (6.5)	$8.2(5.7)^3$	< 0.0001
Oral picture-naming (max 140)	-		-	39.2 (27.6) ⁵	-
Word-triple association (max 70)	_		_	$51.2(10.1)^5$	_
Number calculation task (max 7)	-		-	$6.36(1.1)^5$	-
Lexical decision (max 352)	346.0 (3.7) ¹		$333.2(23.5)^2$	325.3 (23.0) ⁶	0.002
AF, animals/min	23.8 (5.9)	18.2 (3.8)	14.1 (4.2)	$5.6 (4.3)^3$	< 0.0001
VF, verbs/min	21.9 (5.8)	18.2 (5.6)	11.9 (5.0)	$7.0(2.8)^3$	< 0.0001

Table 1
Descriptives and clinical scores. Kruskal-Wallis tests were run to assess group differences of age, education, MMSE, BNT, lexical decision, AF,
and VF. Comparisons between AD and SD of the CDS and GDS scores were performed using the non-parametric Kolmogorov-Smirnov-Test

[†] Normative data are reference values for comparison of the control group (HC) with respect to BNT with N = 32 [81]; AF with N = 94 [82]; VF with N = 67 [83]. ¹n = 16, ²n = 12, ³n = 5, ⁴n = 19, ⁵n = 14, ⁶n = 4. CDS, Cornell Depression Scale; GDS, Global Deterioration Scale; MMSE, Mini-Mental State Examination; BNT, Boston Naming Test; AF, animal fluency; VF, verb fluency.

performed by expert clinicians and was in accordance 218 with the ICD-10 criteria [44]. The patients with AD 219 included in this study underwent a standard clini-220 cal procedure which consisted of examinations such 221 as structural neuroimaging, lumbar puncture, blood 222 analyses, and a neuropsychological assessment (these 223 assessments were part of the clinical routine and 224 only used for diagnosis). Further inclusion criteria for 225 patients from the Stockholm site was a Global Dete-226 rioration Scale lower than 6 (i.e., moderate dementia 227 or milder) and the Cornell Depression Scale below 228 8. Healthy elderly controls were recruited by adver-229 tisement (n = 22) in the Stockholm area. Presence of 230 medical or psychiatric disorders (other than demen-231 tia), intake of drugs affecting the nervous system, or 232 magnetic implants, led to an exclusion from the study. 233 Variables available for all participants included in 234 the study were age, gender, and Mini-Mental State 235 Examination (MMSE). 236

All study participants provided informed consent
prior to the data acquisition. The Shanghai study
was approved by the Institutional Review Board of
the State Key Laboratory of Cognitive Neuroscience
and Learning, Beijing Normal University [33]. The
Stockholm study was approved by the Regional
Ethics Committee of Stockholm, Sweden.

244 MRI data

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MR images were acquired on two sites: The Karolinska Institute in Stockholm, Sweden, and the Huashan Hospital in Shanghai, China.

Stockholm site

MR images were acquired with a 3T Siemens Magnetom Trio scanner (Siemens AG, Erlangen, Germany). Structural images were 3D T1-weighted magnetization-prepared rapid gradient echo (MPRAGE) images using the following parameters: TR = 1900 ms, TE = 2.57 ms, flip angle = 9° , matrix size = 256×256 , field of view = 230×230 mm², slice number = 176 slices, slice thickness = 1 mm, and voxel size = $0.90 \times 0.90 \times 1 \text{ mm}^3$. The structural images were previously used for voxel-based morphometry and published with a different purpose and sample configuration [45, 46]. Functional images were acquired with a 32-channel head coil, using an interleaved EPI sequence (400 volumes; 26 slices; voxel, $3 \times 3 \times 4$ mm³; gap thickness, 0.2 mm; matrix size, 80×80 ; FOV, $240 \times 240 \text{ mm}^2$; TR, 1600 ms; TE, 35 ms).

Shanghai site

Images were acquired with a 3T Siemens MagnetomVerio. Structural images were 3D T1-weighted magnetization-prepared rapid gradient echo (MPRAGE) images using the following parameters: TR = 2300 ms, TE = 2.98 ms, flip angle = 9°, matrix size = 240 × 256, field of view = 240 × 256 mm², slice number = 192 slices, slice thickness = 1 mm, and voxel size = $1 \times 1 \times 1$ mm3. Functional images were acquired with a 32-channel head coil, using an interleaved EPI sequence (200 volumes; 33 slices; voxel, $4 \times 4 \times 4$ mm³; gap thickness, 0 mm; matrix size, 64×64 ; FOV, 256 × 256 mm²; TR, 2000 ms;

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284 Preprocessing of functional MRI scans

We performed pre-processing using SPM12 285 (http://www.fil.ion.ucl.ac.uk/spm). We initially set 286 all images' origin to the anterior commissure, 287 and then performed slice-time correction, realign-288 ment, coregistration, normalization to MNI space 289 $(2 \times 2 \times 2 \text{ mm}^3)$, and smoothing (full width half 290 maximum [FWHM]: 8 mm). Time series data were 291 high-pass filtered (1/128 Hz) and we regressed out 292 14 nuisance parameters (6 movement parameters and 293 their first derivative, white matter, and cerebrospinal 294 fluid). 295

We carefully assessed data quality and inspected 296 the spatio-temporal quality of each scan by compar-297 ing the slow and fast components of the data using 298 DSE (Dvar, Svar&Evar) decomposition [48]. The 299 DSE technique decomposes the dataset into three 300 main components: fast, which is the squared mean 301 difference; slow, which is the squared mean averages, 302 and Evar, which refers to the sum of squares of the 303 two ends of the time series. Subjects with remark-304 ably high divergence (>75%-tile) between Dvar and 305 Svar components were removed, as suggested in Afy-306 ouni & Nichols [48]. Therefore, we removed one SD 307 and three HC datasets from the analysis. We further 308 excluded two AD subjects, as more than 20% of their 309 DVARS data-point were found to be corrupted. The 310 remaining subjects were scrubbed as suggested by 311 Power et al. [49]. Altogether, we excluded six datasets 312 (9.7%) due to poor data quality. We re-run the diag-313 nostics on the final sample and found no difference 314 between groups regarding the DSE diagnostics (one-315 way ANOVA, all p > 0.05). 316

317 Functional connectivity analysis

We investigated FC between each of the 66 tem-318 poral ROIs in three participant groups (AD, SD, 319 and HC). We focused our analysis on the temporal 320 lobes with the following rationale: first, brain regions 321 identified as the origin of atrophy are located in the 322 temporal lobe. Second, a 'crossroad' in FC network 323 disruption in AD and SD was found in the hippocam-324 pus. Third, functional hubs for episodic and semantic 325

memory can be found in the temporal lobe (as outlined above). Fourth, the strongest FC of temporal regions is located within the temporal lobes and concurs with functional networks crucial for language processing, the core clinical feature of SD [50]. The functional parcellation we used is based on restingstate fMRI data which was clustered into spatially coherent regions of homogeneous FC and was evaluated in terms of the generalizability of group level results to the individual [38]. From the 200 ROIs, we used a subset of 66 temporal ROIs that covered at least 5% or more of one of the following temporal structures from WFU Pickatlas 3.0.4 [51]: the superior temporal cortex, the middle temporal cortex, the inferior temporal cortex, the temporal pole, the hippocampus, the parahippocampal cortex, the lingual gyrus, the amygdala, the insular cortex, and the fusiform gyrus; these 66 ROIs are shown in Supplementary Figure 1. Analyzing merely 66 temporal ROIs leads to 2,145 pair wise correlations, which necessitates a strong adjustment for multiple comparisons to control for false positives. Using an even higher number of ROIs, for example comparing 200 ROIs in the whole brain, would require an even stronger correction (correcting for almost 20,000 comparisons). Such corrections would result in a sensitivity too low to detect even substantial FC changes.

We extracted the mean time series from the gray matter (probability > 0.70) of these ROIs to assure that time series were not contaminated with cerebrospinal fluid signals from atrophied areas, resulting in 66 time series per subject. We also assured that time series were not affected by signal dropouts due to dephasing. To address motion and physiological confounds which are global in nature, we applied global signal regression to the time series [52–54]. We created a pair-wise correlation matrix and transformed the correlation coefficient to Z-scores by Fisher's transformation. We conducted a one-way ANOVA for each ROI pair (2,145 tests) to test the null hypothesis of no difference between the three groups. We performed an additional sensitivity analysis with age, mean gray matter density in the temporal cortex, MMSE, and study site as additional covariates. Covariates can be problematic if these differ between groups [55], therefore we report theses sensitivity analyses in the Supplementary Material. From the 2,145 total connections, we found 321 (sensitivity analysis: 324) significant edges that showed a group effect (uncorrected, p < 0.05), and after correcting the *p*-values for multiple comparisons, seven edges 326

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showed a significant group effect (FDR corrected, p < 0.05).

380 Voxel-based morphometry analysis

We additionally performed a VBM analysis to 381 quantify gray matter loss in the patients from the 382 anatomical T1 images. VBM is a voxel-wise com-383 parison of the local amount of gray matter volume 384 between two groups [56]. We performed the follow-385 ing processing steps: spatial registration to MNI space 386 (voxel size: $1.5 \times 1.5 \times 1.5 \text{ mm}^3$) and tissue segmen-387 tation, bias correction of the intensities, smoothing 388 of the GM images with 8 mm FWHM, and mod-389 ulation by scaling with the total volume so that 390 the resulting amount of gray matter in the modu-391 lated images remained the same as in the native 392 images. In other words, this step removed the intro-393 duced bias from the registration of different brain 394 sizes to MNI space. The Stockholm sample was 395 registered using the European brain template, the 306 Shanghai sample with the East Asian brain template 397 and normalized to MNI space. We used the "Compu-398 tational Anatomy Toolbox" CAT12 [57] and SPM12 399 [58] for the VBM analysis. Statistical inference 400 was performed with the "Statistical Non Parametric 401 Mapping" software SnPM13 using non-parametric 402 permutation/randomization two-sample t-tests with 403 a voxel-wise family-wise error correction (FWE) of 404 0.05. We performed two t-tests and compared the HC 405 group versus the SD group, and the HC group versus 406 the AD group. Unlike in the analyses of the functional 407 data where we excluded six datasets, the structural T1 408 scan from all the subjects were used in this analysis. 409 the group sizes were HC with n = 20, AD with n = 18, 410 SD with n = 24. 411

412 **RESULTS**

We first describe the clinical presentation of the 413 patients included in this study (see Table 1 for details). 414 The SD group performed poorer in MMSE than the 415 AD group (Kruskal-Wallis over all groups: H = 29.5, 416 df = 2, p < 0.0001; Kolmogorov-Smirnov group-wise 417 post-hoc tests: HC-AD Z=1.86, p=0.002, HC-SD 418 Z=2.52, p<0.001, AD-SD Z=1.44, p=0.033). Fur-419 thermore, the SD group showed significantly lower 420 scores in the Boston Naming Test (BNT) than the 421 AD group (Kruskal-Wallis over all groups: H = 23.3, 422 df = 2, p < 0.0001; Kolmogorov-Smirnov group-wise 423 post-hoc tests: HC-AD Z=1.85, p=0.002, HC-424 SD Z = 1.97, p = 0.001, AD-SD Z = 1.95, p = 0.001). 425

Within the SD group, we observed that the impaired 426 performance in picture naming (BNT, Stockholm 427 site; oral picture-naming, Shanghai site) were more 428 pronounced than lexical decision (Stockholm site) 429 and word-triple association (Shanghai site), see 430 Table 1. The group differences between SD and AD 431 in MMSE and BNT are common findings given that 432 the BNT is a semantic task and the MMSE relies on 433 language comprehension, as both semantics and lan-434 guage are typically more affected in SD than AD. 435 Finally, our AD group also showed semantic deficits 436 as compared to the healthy control group (based 437 on BNT, animal fluency, and verbal fluency). These 438 behavioral scores mirror the severe semantic memory 439 deficits in patients with SD. Moreover, the normal 440 calculation ability in the majority of our SD group 441 supported the diagnostic features of SD. In contrast, 442 patients with AD showed a comparably mild seman-443 tic memory deficit, which is in accordance with the 444 expectations. MMSE was the only available neu-445 ropsychological test score for all participants from 446 both sites, whereas the remaining tests were site-447 specific and therefore not comparable. 448

Next, we report the gray matter density found in the patient groups, see Fig. 1. In the SD patients (Fig. 1A), we found two clusters of atrophy. The first was located in the left anterior medial temporal cortex, with a peak effect in the left temporal fusiform cortex (peak *t*-score = 14.0, p_{FDR} = 0.0021, df = 42; location at x = -34, y = -3, z = -36; cluster area 80.7 cm³). The second cluster was located in the temporal fusiform cortex of the right hemisphere (peak *t*-score = 10.6, $p_{\text{FDR}} = 0.0021$, df = 42; location at x = 34, y = -3, z = -34; cluster area 40.1 cm³). In the AD patients, we found two clusters with lower GM volume compared to controls in the left amygdala (peak *t*-score = 8.72, $p_{\text{FDR}} = 0.006$, df = 36; location at x = -26, y = -10, z = -12; cluster area 7.23 cm³) and the right amygdala (peak *t*-score = 7.49, $p_{FDR} = 0.006$, df = 36; location at x = 22, y = -3, z = -15; cluster area 7.47 cm³), see Fig. 1B. A commonly expected hippocampal atrophy was yielded only with a more liberal threshold (Supplementary Figure 2).

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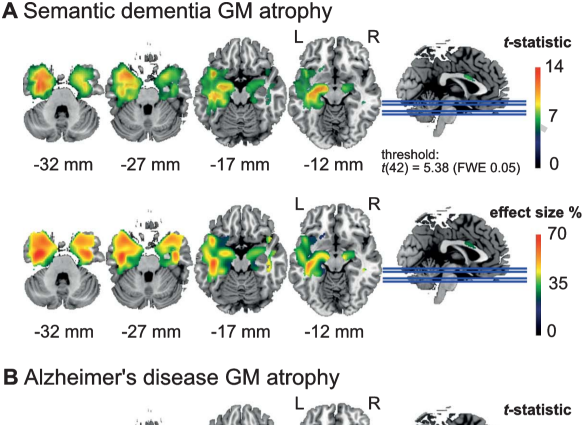
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To achieve the main goal of this study, we analyzed the functional connectivity of 56 participants using 66 functional ROIs of the temporal cortex and related sub cortical areas (see complete correlational matrix in Supplementary Figure 3). Seven connections (FC between ROI pairs) demonstrated a significant difference between the three groups after correcting for multiple comparisons (FDR corrected, p < 0.05). A detailed characterization and test statistics of these



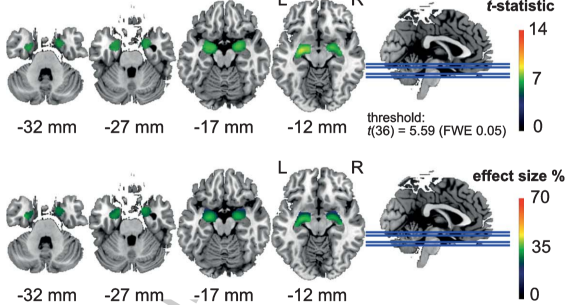


Fig. 1. Areas with significantly lower (voxel-level) gray matter (GM) density (top) and effect size in terms of percentage GM reduction (bottom) in (A) the semantic dementia (SD) patients (n = 24) and (B) Alzheimer's disease (AD) patients (n = 18) compared to the healthy elderly control group (n = 20). SD patients showed reduced GM density in widespread areas of the left anterior temporal cortex including the temporal pole, while the AD patients showed reduced GM density in the amygdala. SD patients showed more severe GM loss with up to 70% reduction, and AD patients with up to 40% reduction in some areas.

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Edge no	. ROI no.	ROI no.	Region	Region	F	FDR adj. p
1	129	11	Left anterior superior temporal gyrus/middle temporal gyrus/insular cortex	Left posterior middle temporal gyrus/superior temporal gyrus	10.86	0.034
2	85	24	Right lateral inferior occipital cortex/lateral superior occipital cortex	Left posterior superior temporal gyrus/central opercular cortex/parietal opercular cortex/planum temporale	11.52	0.030
3	198	32	Right fusiform cortex/parahippocampal gyrus	Right inferior temporal pole	12.64	0.026
4	70	37	Left lingual gyrus/intracalcarine cortex/precuneus cortex	Left posterior hippocampus/thalamus	13.18	0.026
5	89	37	Right lingual gyrus/intracalcarine cortex	Left posterior hippocampus/thalamus	10.95	0.034
6	153	71	Right anterior middle temporal gyrus/superior temporal gyrus	Right orbitofrontal cortex	12.42	0.026
7	112	72	Left orbitofrontal cortex/insular cortex	Left anterior inferior temporal gyrus/middle temporal gyrus	12.06	0.026
	edge no. 1 L_asTmp		edge no. 2 p R_liOcc ↔ L_psTmp	edge no. 3 R_Fusi ↔ R_iTmpPo	edge no. 4 L_Ling ↔	
1. 1. N ^{0.} 0. -0.	0 5 0		1.5 1.0 N 0.5 0.0 -0.5 SD AD HC	1.5 1.0 0.5 0.0 -0.5 SD AD HC 1.5 1.0 0.5 0.0 -0.5 SD AD HC	SD A	D HC
1. 1. N 0. 0. -0.	0 5 0	L_Hipp	edge no. 6 R_mTmp ↔ R_FrtOrb 1.5 1.0 0.5 0.0 -0.5 SD AD HC	edge no. 7 L_FrtOrb ↔ L_aiTmp 1.5 1.0 0.5 0.0 -0.5 SD AD HC		

Table 2 Seven functional connections that demonstrated significant group differences

Fig. 2. Z-scores of seven connections (edges 1-7) with significant group differences. Post-hoc tests between the three groups were performed, and significant group differences are denoted with red horizontal lines (see Table 2 for a detailed description of the ROIs). Ring-shaped circles represent single subject data points. Filled circles represent outliers.

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Post-hoc Tukey HSD p-values for the three single comparisons (rows) and for each of the seven ROI-pairs that had a significant group effect (columns). Significant values reflect that the group effect was driven by a specific group level contrast

Edge No.	1	2	3	4	5	6	7
AD versus HC	0.098	0.082	0.98	0.54	0.22	0.0005	1.00
SD versus HC	< 0.0001	0.044	0.0002	< 0.0001	< 0.0001	< 0.0001	0.0004
SD versus AD	0.064	< 0.0001	0.0004	0.002	0.024	0.97	0.0004

seven connections are shown in Table 2; the Z-values 478

for the significant connections are depicted in Fig. 2.

479 We performed post-hoc tests (Tukey HSD) for sin-480

gle comparisons of the three groups to investigate the particular group contrasts that drove the significant group effect (Table 3). We found that most differ-

ences were related to the SD patients with significant 484 changes in all of the seven connections. SD patients 485 showed lower FC in 6 out of 7 connections compared 486 to HC, and higher FC in one connection (edge no. 2) 487 compared to HC. This higher FC in the SD patients 488 was also significantly higher compared to the AD 489 patients. The AD patients had a lower FC compared 490 to HC patients in only 1 out of the 7 connections 491 (edge 6). Comparing the two patient groups SD, ver-492 sus AD, we found that SD had a significant lower FC 493 in 4 connections (edges 3, 4, 5, 7). 494

We visualized the connectivity structure and con-495 nection strengths, see Fig. 3. The SD patients 496 generally had a much lower connectivity compared to 497 the other two groups. An exception was the stronger 498 contra lateral connection between the right lateral 499 inferior occipital cortex and the left posterior supe-500 rior temporal gyrus (edge no. 2). The AD patients 501 showed a lower FC compared to HC between the right 502 middle temporal gyrus and the right frontal orbital 503 cortex (edge no. 6). A common finding in all the three 504 groups was that the FC between the right fusiform 505 cortex and the right inferior temporal pole was the 506 strongest (no. 3). 507

The sensitivity analysis with age, mean gray mat-508 ter density in the temporal cortex, MMSE, and study 509 site as covariates produced statistically significant 510 differences in the same seven edges as reported 511 above; however, the assumption of the ANCOVA, 512 the independence between the patient group and the 513 covariates was not met. For results of the sensi-514 tivity analysis with covariates, see Supplementary 515 Tables 1-4. 516

517 DISCUSSION

In this study, we compared functional connectiv-518 ity between SD, AD, and HC using a functional 519 parcellation of 66 ROIs of the temporal cortex and 520 hippocampus to investigate intra-temporal connec-521 tions and connections with contra lateral temporal 522 regions. The overall picture that emerges is that 523 between the majority of the significant ROIs, SD 524 demonstrated the most striking decrease in FC. In 525 the AD group, most differences compared to the HC 526 group did not reach significance. We believe that the 527 often described disconnections found in AD were not 528 detected in our study due to the mild progression of 529 the disease in our AD group. One reason could be that 530 the remaining gray matter volume in brain regions 531 typically affected by neuronal degeneration was suffi-532

cient to maintain an intact FC to remote areas. In other words, the damage found in mild stages might affect the intra-regional processing in local neuronal populations, whereas the inter-regional (i.e., network) FC would be affected during more advanced AD progression [59]. Future studies will require larger sample sizes to demonstrate smaller changes in FC seen even with mild state impairments.

The most intriguing finding of our study for the SD group was the decreased FC between the left posterior hippocampus and left/ right lingual gyri (edges 4 and 5). These disruptions are characteristic for the neurophysiological basis of the SD patients' typical symptomatology involving an impaired semantic memory. For instance, Sormaz et al. [60] recently showed a correlation of FC between left the hippocampus and the lingual gyrus with topographic memory, and a correlation of semantic memory performance with FC to the intracalcarine cortex, a finding consistent with our results.

Functional connectivity between the left anterior superior/middle temporal gyrus/insula and the left posterior middle/superior temporal gyrus was decreased in SD compared to HC (edge no. 1). It is important to note that this is the single connection that showed an FC difference between SD and HC exclusively (i.e., a finding specific for the SD-HC group single-comparison while neither AD-HC nor AD-SD were significant). These regions are commonly associated with cross-modal integration (as is the hypothesized semantic hub) of auditory and language processing, as well as the processing of the emotionally relevant context. Hence, this finding might reflect the severe semantic deficits in SD (see Table 1) that are manifested by the loss of conceptual knowledge [7].

The single connection that showed increased FC in SD compared to the other groups (edge no. 2) was between the left posterior superior temporal gyrus/parietal opercular cortex/planum temporaleand the right lateral inferior/superior occipital cortex. The temporal brain areas that constitute this connection are important for early context integration of acoustically presented words [61], lexico-semantic retrieval [62], and are part of a supramodal semantic network [63]. The occipital ROI of this connection sub serves visual integration. Thus, an increased FC between these regions might reflect a functional reorganization that is characterized by supporting language comprehension using more sensory inputs. Moreover, this result indicated a reduced hemispheric functional specialization and perhaps an attempt to pool

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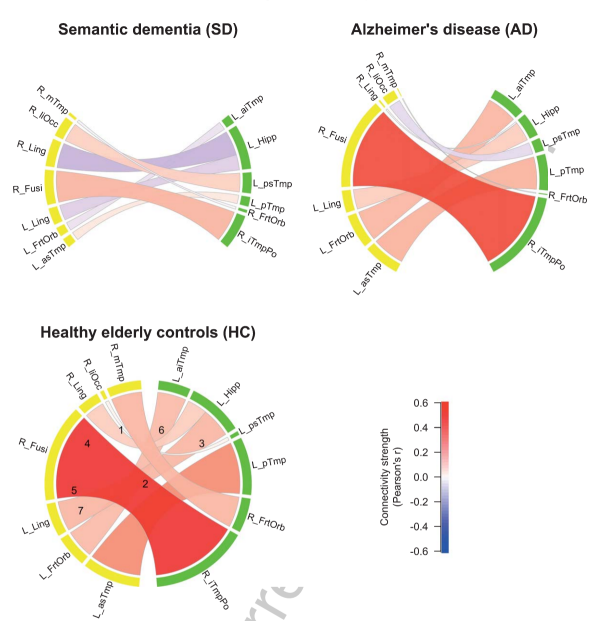


Fig. 3. Functional connectivity (FC) strengths of the three groups. Color shades and thickness of the links are proportional to FC strengths; shades of red reflect positive, shades of blue negative strengths. Numbers in HC group indicate edge numbers (see Table 2 for a detailed description of the ROIs). ROIs in the left hemisphere are labeled yellow, ROIs in the right hemisphere labeled green.

resources that are spared by the pathological developments in SD.

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In comparison with AD and HC, SD patients showed a lower FC between the functional ROI encompassing the right fusiform/parahippocampal gyri and the right inferior temporal pole. Disruption of this connection (no. 3) can be viewed as SD-typical, as the functional profile of the involved regions conforms to SD symptomatology. In particular, the right temporal pole is crucial for non-verbal (e.g., visual) object recognition, which is a hallmark impairment in SD associated with the loss of semantic knowledge [64, 65]. The right fusiform gyrus on the other hand is associated with working memory for faces, face perception, and non-verbal associative semantic knowledge [66–68], and the right parahippocampal gyrus is associated with working memory for object location as well as a function as an episodic buffer [69, 70]. In line with this, the patients with SD in the present study showed severe object recognition

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deficits assessed with BNT and oral picture-naming
 (Table 1), even though no behavioral data about face
 perception or non-verbal semantic knowledge was
 available.

Similar to connection no. 3, FC was reduced 609 in SD compared with AD and HC between the 610 ROI comprising left lingual/intracalcarine/precuneus 611 cortex and the ROI including left posterior hippocam-612 pus/thalamus (connection no. 4). This finding is in 613 line with Seeley et al. [14], who reported the medial 614 temporal lobe as part of an SD-vulnerable network. 615 Thus, in addition we showed a possible contribu-616 tion of the primary visual (intracalcarine cortex), 617 visual memory (lingual gyrus) and self-awareness 618 (precuneus, i.e., default mode network) regions to that 619 semantic network. It might appear surprising that the 620 FC of the AD group was not significantly reduced in 621 this connection, despite the commonly known medial 622 temporal lobe atrophy and the pivotal role of the hip-623 pocampus in episodic memory encoding [36, 71]. 624 However, functional and anatomical changes do not 625 necessarily overlap, and for instance, stable FC of the 626 left hippocampus in early AD (except with right lat-627 eral prefrontal cortex) has been reported previously 628 [29]. 629

Lower FC in SD than in AD (and HC) was 630 also found between the right lingual/intracalcarine 631 cortex and the left posterior hippocampus/thalamus 632 (connection no. 5). Therefore, connections between 633 bilateral lingual gyri and the left hippocampus were 634 detected in our HC sample (for illustration, see Fig. 3, 635 connections no. 4 and 5), whereas either of them were 636 damaged in SD, but not in AD. This supports the 637 recent indication of a hippocampal contribution to 638 the semantic memory network [36]. Because episodic 639 memory is relatively spared in SD, the connections 640 between the left posterior hippocampus and the bilat-641 eral lingual gyri might contribute to the semantic 642 memory network. On the other hand, we did not find 643 an expected decrease of FC in connection no. 4 in AD, 644 although the precuneus and hippocampus contribute 645 to episodic memory, which is typically impaired in 646 AD. However, we have to bear in mind that our analy-647 sis was restricted to temporal lobe FC and thus did not 648 cover the entire episodic memory network, including 649 brain regions located in frontal and parietal lobes. In 650 addition, no episodic memory data were available for 651 the entire sample of our study. Future studies should 652 investigate additional ROIs from the aforementioned 653 areas using larger sample sizes to tackle the increased 654 number of connections and multiple testing correc-655 tions that are associated with larger networks. 656

The only FC reduction common to both SD and AD compared with HC was found in connection no. 6. The functional role of the involved regions suggests an association with a frequently observed clinical presentation of AD and SD characterized by apathy and agitation, associated with the right orbitofrontal cortex [72, 73], and impairments in social behavior related to the right anterior temporal lobe [74]. According to Olson et al. [75], social knowledge is part of semantic memory and involves memory about people including biographical information. Nonetheless, caution is advised with comparing social or semantic deficits between AD and SD: both symptoms have different onsets or severities within disease stages, as well as different characteristics. Furthermore, we did not have data on social behavior or apathy/agitation of our patients. Regardless, we added a common pathway to the crossroad described by La Joie et al. [36]. They suggested that the hippocampus is a converging hub of an (AD-affected) episodic and a (SD-affected) semantic network. Accordingly, our data indicated that besides a shared damaged hub in AD and SD, the functional connection between the right anterior middle/ superior temporal gyri and the right orbitofrontal cortex might be a second candidate for the neuropathology shared in both clinical populations.

The final significant connection (no. 7) of the present study was found between the left orbitofrontal cortex and the left anterior inferior and middle temporal gyri. The literature suggests a functional role of this connection in deficient socio emotional abilities that are found predominantly in the behavioral variant of FTD [76], and in higher level object representation, involving language and auditory processing. Unlike in connection no. 6, the AD group did not show an impaired FC of the orbitofrontal regions with the ipsilateral temporal cortex. Thus, one might speculate about a bilateral breakdown of orbitofrontal to temporal connections in SD, which might be related to the severity of the semantic deficit.

This study entailed a number of study design limitations that need to be taken into account while interpreting the results. Even though the overall sample size is large, the sample sizes of the three subgroups are considered small (16–23 individuals). Larger studies need to be conducted, however, this is especially challenging for SD given its low prevalence. Therefore, we pooled two SD samples from two different sites with different scanners. However, most individuals of the SD group and none of the AD and HC groups were from the Shanghai site, 657

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which is a violation of acknowledged study design 709 standards and a potential confound. Moreover, the 710 diagnostic criteria of the two sites for SD were not 711 identical. The data from different MRI sites may have 712 different noise levels such as thermal noise, physio-713 logical noise, and motion [52, 77]. These artifacts are 714 often global in nature, and global signal regression 715 (GSR) can successfully remove these and standard-716 ize the data between sites and across individuals. GSR 717 can introduce negative correlations; however, GSR 718 can also improve the specificity of positive corre-719 lations [78]. Importantly, in this study, we do not 720 interpret absolute negative correlations and solely 721 compare relative differences in correlations between 722 groups. We conducted a sensitivity analysis with the 723 study site as covariate which yielded the same results. 724 However, assumptions of independence between the 725 covariates and the patient groups were not met. The 726 present limitation of the unbalanced study design can-727 not entirely be removed by an analysis of covariance. 728 Thus, future studies should measure different patient 729 populations across different scanner sites and ideally 730 achieve balanced groups across sites. Harmoniza-731 tion techniques [79] can further improve data quality. 732 However, the application of harmonization methods 733 in unbalanced groups is questionable as these not 734 only eliminate scanner effects, but also the effects of 735 interest [80]. Moreover, interpretation of group differ-736 ences between AD and SD should take into account 737 that the two dementia groups were not matched for 738 disease stage (i.e., SD showed more severe deficits 739 than AD). The sensitivity analysis with GM density 740 as covariate was in line with our results. Likewise, 741 more symptom specific behavioral scores (other than 742 MMSE) could have aided an in-depth interpretation 743 of altered FC edges in the patient groups. Lastly, our 744 analysis did not cover all brain regions potentially rel-745 evant for AD and SD. However, the choice to limit the 746 scope to the temporal lobe has three reasons: first, the 747 distinct temporal lobe atrophy is crucial for AD and 748 SD differentiation. Second, the temporal lobe is piv-749 otal in both semantic and episodic memory functions. 750 Third, the definition of ROIs within the temporal lobe, 751 even though using an arbitrary selection threshold of 752 5% (or more) of overlap of the ROIs with any tem-753 poral structure, may be altogether less arbitrary and 754 biased compared to subjectively selecting ROIs based 755 on expectations and literature. 756

To summarize the main findings of our study,
the cohort of patients with SD yielded a number of
distinct ipsilateral and contra lateral connections of
the temporal lobe that showed a significant reduc-

tion in FC. These connections included the regions on which our predictions were based on (i.e., hippocampus, fusiform gyrus, and temporal pole). Two functional connections were intriguing due to their distinctiveness from the other groups: the first was the connectivity breakdown between left posterior hippocampus and bilateral lingual gyri, likely reflecting the neuronal underpinning of semantic memory loss. Second, a bilateral disruption of connectivity between temporal and frontal lobes was found. This aligns well with the pathophysiology within the FTD spectrum and especially with SD.

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FC in AD was relatively intact compared to SD, which contradicted our hypothesis. The only connection with significantly reduced FC encompassed the right orbitofrontal cortex and the right anterior temporal lobe (no. 6), which we identified as an AD/SD-common pathway. Additionally, Fig. 2 illustrates that our AD group had a lower FC than the HC in connections no. 1, 2, and to a smaller extent no. 5 which all missed significance. These FC signatures in the AD group could be attributed to their mild stage of symptom progression (MMSE of 24.5), and potentially an early marker of the disease, but larger and longitudinal studies are needed.

Following the "cortically distributed plus semantic hub" theory, several connections were found to be significantly altered in the present study, which affected the anterior temporal lobe - semantic hub regions (no. 1, 3, 6, and 7). Moreover, their counterparts were partly localized in the modality-specific regions described by Patterson et al. [5], but also in orbitofrontal regions. This agreement of our results with the arguments in Patterson et al. [5] supported the "distributed plus hub" theory, because we found altered FC in connections between the hub and the modality-specific regions. Taken together, this study presents an alternative concept to investigate the understanding of distinct pathophysiological changes in AD and SD that are related to disruptions of functional networks in the temporal lobe. The unique aspect of our study was the definition of ROIs based on functional brain segregation rather than anatomy for FC analysis. Due to the comparably strict statistical approach and the predefined choice of ROIs, our study provided a fine-grained overview of FC aberration related to temporal lobe function in AD and SD. However, comparability was limited owing to different study sites using partially different diagnostic criteria and data acquisition procedures. We emphasize here that this was an exploratory study with the motivation of gathering MRI data of a rare

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condition (SD) from two study sites to increase sta-813 tistical power. The downside of this approach was 814 that the retrospective characteristic caused unbal-815 anced recording of the study groups across the two 816 MRI scanning sites. This required the conduction 817 of several control analyses to mitigate the occur-818 rence of false contrasts between the groups. Thus, 819 our findings ideally motivate future studies for repli-820 cation with harmonized MRI acquisition parameters 821 and balanced subject numbers between study sites, 822 concurring with an optimal research practice. 823

824 DATA AVAILABILITY

Raw imaging data can be requested from the corresponding author. Aggregated data and analysis scripts to generate all results and figures are available at OSF (https://osf.io/t4jnv/).

829 ACKNOWLEDGMENTS

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839 SUPPLEMENTARY MATERIAL

The supplementary material is available in the electronic version of this article: https://dx.doi.org/ 10.3233/JAD-191113.

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