

Investigating Compensatory Brain Activity in Older Adults with Subjective Cognitive Decline

Christine Krebs^{a,b,*}, Esther Brill^{a,b,c}, Lora Minkova^a, Andrea Federspiel^d, Frauke Kellner-Weldon^e,
Patric Wyss^a, Charlotte E. Teunissen^f, Argonde C. van Harten^g, Anna Seydell-Greenwald^h,
Katharina Klink^a, Marc A. Züst^a, Anna-Katharine Brem^{a,i} and Stefan Klöppel^a

^aUniversity Hospital of Old Age Psychiatry and Psychotherapy, University of Bern, Bern, Switzerland

^bSwiss Institute for Translational and Entrepreneurial Medicine, Bern, Bern, Switzerland

^cGraduate School for Health Sciences, University of Bern, Bern, Switzerland

^dTranslational Research Center, University Hospital of Psychiatry and Psychotherapy, University of Bern, Bern, Switzerland

^eSection Neuroradiology of the Department of Radiology, Cantonal Hospital Lucerne, Lucerne, Switzerland

^fNeurochemistry Laboratory, Department of Clinical Chemistry, Amsterdam Vrije University, Amsterdam, Netherlands

^gAlzheimer Center Amsterdam, Department of Neurology, Amsterdam Neuroscience, Vrije Universiteit Amsterdam, Amsterdam, the Netherlands

^hCenter for Brain Plasticity and Recovery, Georgetown University, Washington, DC, USA

ⁱDepartment of Old Age Psychiatry, Institute of Psychiatry, Psychology and Neuroscience, King's College London, London, United Kingdom

Handling Associate Editor: Jodie R Gawryluk

Accepted 16 February 2023

Pre-press 21 March 2023

Abstract.

Background: Preclinical Alzheimer's disease (AD) is one possible cause of subjective cognitive decline (SCD). Normal task performance despite ongoing neurodegeneration is typically considered as neuronal compensation, which is reflected by greater neuronal activity. Compensatory brain activity has been observed in frontal as well as parietal regions in SCD, but data are scarce, especially outside the memory domain.

Objective: To investigate potential compensatory activity in SCD. Such compensatory activity is particularly expected in participants where blood-based biomarkers indicated amyloid positivity as this implies preclinical AD.

Methods: 52 participants with SCD (mean age: 71.00 ± 5.70) underwent structural and functional neuroimaging (fMRI), targeting episodic memory and spatial abilities, and a neuropsychological assessment. The estimation of amyloid positivity was based on plasma amyloid- β and phosphorylated tau (pTau181) measures.

Results: Our fMRI analyses of the spatial abilities task did not indicate compensation, with only three voxels exceeding an uncorrected threshold at $p < 0.001$. This finding was not replicated in a subset of 23 biomarker positive individuals.

*Correspondence to: Christine Krebs, PhD, University Hospital of Old Age Psychiatry and Psychotherapy, Bolligenstrasse 111, 3000 Bern 60, Switzerland. Tel.: +41(0)31 932 88 27; E-mail: christine.krebs@upd.unibe.ch.

Conclusion: Our results do not provide conclusive evidence for compensatory brain activity in SCD. It is possible that neuronal compensation does not manifest at such an early stage as SCD. Alternatively, it is possible that our sample size was too small or that compensatory activity may be too heterogeneous to be detected by group-level statistics. Interventions based on the individual fMRI signal should therefore be explored.

Keywords: Blood-based biomarkers, episodic memory, functional MRI, neuronal compensation, spatial abilities, subjective cognitive decline

INTRODUCTION

Subjective cognitive decline (SCD) is defined as self-perceived cognitive decline that does not manifest in cognitive tests and affects around 20% of individuals older than 65 years [1]. SCD might have different causes (e.g., neurodegenerative disorders, psychiatric disorders, or head trauma) [2]. Despite achieving normal results in neuropsychological tests, the average performance in SCD is somewhat lower in episodic memory measures of immediate and delayed verbal recall, compared to healthy older adults [3, 4]. Regardless of variations in the trajectory of SCD, it is considered an at-risk state for the development of Alzheimer's disease (AD), particularly when the individual is worried about the decline. Longitudinal studies show that SCD approximately doubles the risk of developing manifest dementia in the following years within five years [5]. This is in line with several studies that reported increased biomarkers of AD in SCD [6]. Studies investigating amyloid- β for example, reported greater levels of amyloid depositions in positron emission tomography (PET) in participants with greater SCD [7, 8]. Therefore, increased amyloid levels might be associated with SCD many years before cognitive impairment manifests [6]. The recent development of amyloid- β blood biomarkers allows the estimation of PET amyloid positivity throughout the different phases of the AD spectrum [9]. Tau depositions are another biomarker of AD but also SCD [6]. pTau181 has been associated with amyloid and tau PET positivity in healthy older adults and participants with AD [10]. These blood-based biomarkers have the potential to identify amyloid positive individuals, which might be in a preclinical stage of AD (SCD likely due to AD).

There seems to be a complex interaction between brain activity, brain area and disease stage in cognitive impairment. In the memory domain, a recent study in individuals with SCD, mild cognitive impairment (MCI) and healthy controls reported an inverse quadratic relationship for task-related activation in

an associative memory task in the left parietal lobe across the different study samples. For the hippocampus and other brain regions a negative linear relationship described the activation better than the quadratic function in relation to the volume, i.e., activation was constantly greater when volume became lower. The SCD classification included APOE genotyping ($\epsilon 4$ carrier) and/or biomarker evidence (left or right hippocampal volume one standard deviation below the mean of the healthy control sample) to increase the likelihood of including participants with preclinical AD [11]. In another study, hyperactivity in the hippocampus at baseline predicted increased longitudinal amyloid deposition, which was not the case for cortical regions. This indicates that hippocampal hyperactivity is related to pathological effects, while greater cortical activity reflects compensatory processes [12]. Evidence for compensatory activity in SCD has also been found outside the parietal cortex. In individuals with subjective memory complaints, the right dorsolateral prefrontal cortex showed greater activity during cued recall in a face-occupation task, while task performance in this sample was not significantly different from healthy controls. The face-occupation task assesses associative episodic memory and especially the recall condition is a sensitive test for early memory impairment in the course of AD [13]. Another study reported greater prefrontal activity in participants with subjective memory decline in an episodic memory encoding task as an indicator of compensatory activity [14].

In addition to the results from functional neuroimaging, two systematic reviews summarizing structural brain changes in SCD reported heterogeneous results. However, a lower hippocampal volume has been a consistent finding [6, 15]. The hippocampus is part of the medial temporal lobe (MTL), which is affected by neuropathology (e.g., atrophy) early in AD whereas neocortical structures like the parietal lobe are affected later [16]. Therefore, many imaging studies have focused on the MTL. The parietal lobe is less investigated in SCD even though it is prone to exhibit early amyloid deposition [17, 18]. Both

120 the MTL as well as the parietal lobe are relevant in
121 episodic memory [19–22].

122 In the spatial domain, impairments are observed
123 early in the course of AD. Delayed recall performance
124 in visuospatial memory tasks like the delayed recall
125 in the Rey–Osterrieth complex figure test has shown
126 the ability to discriminate MCI from healthy aging
127 [23]. Spatial abilities include different subfunctions
128 like spatial perception and mental rotation [24]. A
129 meta-analysis reported consistent activity in the supe-
130 rior and inferior parietal lobes in both hemispheres in
131 mental rotation and spatial imagery tasks in healthy
132 subjects [25]. While no study with functional mag-
133 netic resonance imaging (fMRI) investigated spatial
134 abilities in SCD, greater activity in parietal and
135 temporal areas has been reported in MCI. As no
136 significant difference in task performance between
137 the MCI sample and healthy controls was found this
138 might indicate compensatory activity [26].

139 Overall, there is evidence for compensatory activ-
140 ity in SCD, but data is scarce. More data for
141 this population is especially important because
142 SCD allows the study of early compensatory brain
143 changes. But there exists no clear definition of how
144 neuronal compensation can be addressed. Gregory
145 et al. [27] defined a model to operationalize com-
146 pensatory brain activity in neurodegeneration due to
147 Huntington’s disease. This model includes three vari-
148 ables: task performance, a proxy of disease severity,
149 and brain activity. In an early disease-stage accom-
150 panied by normal performance, compensation through
151 greater brain activity is possible. Generally, com-
152 pensatory activity is seen in early or mild cases
153 of neurodegeneration. In later disease-stages per-
154 formance starts to decline as disease severity increases
155 and compensation is no longer possible [27]. In accor-
156 dance with previous studies, we hypothesized that
157 compensatory effects in SCD will be present in pari-
158 etal [11] and/or frontal [13, 14] brain regions and
159 will be most pronounced in subjects with probable
160 amyloid positivity.

161 The aim of this study was to investigate the exist-
162 ence of compensatory brain activity in SCD. As
163 a lower hippocampal volume was a stable finding
164 in studies investigating SCD, we used hippocampal
165 atrophy as an indicator of potential neurodegenera-
166 tion associated with SCD. We deployed the model
167 of Gregory et al. [27] in fMRI tasks on episodic
168 memory and spatial abilities. We selected these tasks
169 based on their involvement of the parietal lobe and the
170 early impact of AD related pathologies on task per-
171 formance. In the episodic memory task we focused

172 on the cued recall condition as this is a sensitive test
173 for early memory impairment in AD [13]. We defined
174 successful neuronal compensation as greater activity
175 in brain regions correlated with greater task perfor-
176 mance in the presence of high hippocampal atrophy.
177 Hippocampal atrophy served as a marker of disease
178 related neuropathology. Compensatory effects might
179 be most pronounced in SCD if it is a preclinical state
180 of AD. Therefore, we repeated the fMRI analyses in a
181 subsample with probable amyloid positivity accord-
182 ing to blood biomarkers for amyloid- β or pTau.

183 METHODS

184 This bi-centric study (Bern and Lucerne;
185 Switzerland) was approved by both local Ethics
186 Committees and registered on ClinicalTrials.gov
187 (NCT04452864). All study participants gave
188 informed consent before the first study visit.

189 *Participants*

190 52 Caucasian participants (mean age 71.00, SD:
191 5.70, mean years of education: 15.15, SD: 3.06) were
192 included in the analyses. The data were collected
193 as part of the baseline assessment of a cognitive
194 training study [28]. This assessment was the first
195 in person contact of the participants with a member
196 of the research team, therefore, we do not assume
197 any effect of the planned intervention or general
198 study setting on the results. We included partici-
199 pants who reported subjective cognitive complaints
200 and related worries as this increased the probability
201 to include participants with SCD likely due to AD
202 [2]. To identify SCD, participants completed a ques-
203 tionnaire on memory related concerns and attentional
204 deficits during the last 12 months. Language or other
205 cognitive abilities were not assessed. This ques-
206 tionnaire is based on suggested criteria for SCD [29]. We
207 included participants only if they reported a decline in
208 memory or attention functions and expressed related
209 worries. For this categorical questionnaire no cut-off
210 scores are available.

211 Other inclusion criteria were age between 60–85
212 years, native or fluent German speakers and nor-
213 mal or corrected to normal vision. Exclusion criteria
214 included a diagnosis of cognitive impairment (MCI or
215 dementia), a severe neurological or acute psychiatric
216 disease, substance abuse, current psychoactive medi-
217 cation, contraindication for MRI (i.e., metal implants)
218 or stroke in previous history. The diagnosis of MCI
219 was based on established criteria [30]. Furthermore,

participants which scored below 23 points in the MoCA were excluded from this study as this is an indicator of objective cognitive decline [31].

For 21 participants no increased cardio-vascular risk factors were identified (i.e., no self-reported high blood pressure, cardiac disease, or abnormalities in the MRI data). 18 participants reported high blood pressure and 5 participants reported heart problems (e.g., heart attack or auricular fibrillation in the past). T1 weighted MRI images were investigated by a clinical neuroradiologist to rate hypointensities, which could reflect a vascular component explaining cognitive impairment. 34 participants of our sample had a Fazekas score (i.e., presence of white matter lesions) [32] of 0, for 10 participants the score was 1, for 7 participants the score was 2 and for 1 participants the score was 3. Of note, we did not acquire FLAIR or similar sequences, which would have been more sensitive measure of white matter lesions.

51 participants were included in the fMRI model for the episodic memory fMRI task, one participant was excluded due to motion artefacts.

Blood samples from 38 participants were available. To investigate neuronal compensation specifically in SCD likely due to AD, we repeated all analyses with a subsample with blood biomarkers indicating amyloid positivity ($n = 23$). In this subsample, one participant had to be excluded from analyses in the episodic memory fMRI task due to motion artefacts.

Neuropsychological assessment and behavioral composite scores

The neuropsychological test battery included the Montreal Cognitive Assessment (MoCA) [33], auditory verbal learning test (AVLT) [34], Rey–Osterrieth complex figure (RCF) [32], flanker test [36], graded naming test [37], semantic fluency, digit span forward and backward, and questionnaires related to the cognitive training intervention. Additional questionnaires assessed situational motivation [38], quality of life [39], activities of daily living [40], handedness [41] and depressive symptoms [42]. Furthermore, we assessed subjective cognitive complaints in self and informant rated versions. The MoCA was the only test performed as paper-pencil version, all other tests and the questionnaires were administered using a tablet (iPad, 7. Generation) with the Apollo App [43].

We computed a composite score of episodic memory and spatial abilities based on raw test scores. The episodic memory (memory) composite score included the learning sum, immediate and delayed

recall of the AVLT. The spatial abilities composite score included encoding, immediate and delayed recall from the RCF. Before calculating the behavioral composite scores, two principal component analyses (PCAs) were performed, one for the three AVLT and the other for the RCF scores. In a next step, we centered and standardized the three test scores included in the composite score. If the PCA showed different loadings for the test scores (i.e., the differences between two loadings were larger than 0.05) within one composite score, the centered and standardized scores were weighted according to their loading and a mean score was calculated, resulting in one memory and one spatial abilities composite score.

Blood-based biomarkers

We used the amyloid- $\beta_{42/40}$ ($A\beta_{42/40}$) ratio and pTau181 measures to identify participants with probable AD-specific neuropathologies and repeated the fMRI analyses with a subsample with positive blood biomarkers for amyloid positivity.

For blood biomarker measurement in this study we used N4PE Simoa immunoassays (IA-N4PE) developed by Amsterdam University Medical Center, Amsterdam, the Netherlands, and ADx Neurosciences, Ghent, Belgium, and commercially available from Quanterix, Billerica, Massachusetts [44]. Cut off scores of 0.06 for the $A\beta_{42/40}$ ratio and 1.8 pg/ml for pTau181 were used. These are based on unpublished data in which 1111 participants with known amyloid status based on CSF or amyloid PET from the Amsterdam Dementia Cohort were analyzed (AUC for $A\beta_{42/40}$ 0.735 and for p-tau181 0.828).

Study procedures

The duration of the study visit was approximately 3 h, including the MRI session. After signing the consent form, the MoCA and tablet-based cognitive tests were performed. Next, participants practiced all conditions of the fMRI tasks outside the MRI scanner to ensure task comprehension. The MRI session itself took around 50 min. The session started with a resting-state fMRI (rs-fMRI) while fixating a cross, followed by the face-occupation task (episodic memory) during task-based fMRI, T1 w structural imaging, a visual construction task targeting spatial abilities during task-based fMRI, and an arterial spin labelling (ASL) protocol. This listing of MRI sequences corresponds to the order in MRI

318 data acquisition. Data on rs-fMRI and ASL are not
319 reported here.

320 *Face-occupation task*

321 Episodic memory was assessed with a blocked
322 face-occupation task (duration: 638 s, Fig. 1), which
323 was based on a previous study [45]. This paradigm
324 induces activity in the MTL and occipital brain areas
325 [46]. Additionally, cued recall tasks have been sug-
326 gested as sensitive markers for AD [13, 47] and have
327 been related to CSF markers of AD [47].

328 The task included four conditions which were pre-
329 sented in the same order (encoding, cued recall,
330 recognition, control condition) in six runs. Before
331 the encoding block a task instruction was shown
332 (“Please remember the people and their job. Does the
333 face fit to the job? Yes = 2, No = 3”). This text disap-
334 peared after 4 s and the first of five face-occupation
335 associations was shown. Below the picture and the
336 occupation the question “Does the face fit to the job?
337 Yes = 2, No = 3” was displayed. The aim of this ques-
338 tion was to induce a deeper level of encoding [13].
339 This block was followed by the control condition.
340 The instruction text appeared again for 4 s (“Sil-
341 houettes: Male or female? 2 = Female, 3 = Male”).
342 This text was followed by five head contours with
343 the question “2 = Female/3 = Male” right aside. Then
344 the instruction for the cued recall condition was dis-
345 played for 4 s (“What is this person’s education?
346 2 = University degree, 3 = Apprenticeship”). During
347 the cued recall condition, a previously learned face
348 appeared as cue with the text “2 = University degree/
349 3 = Apprenticeship” right aside. For the recognition
350 condition the instruction was “What is this person’s
351 occupation?”. This instruction screen was followed
352 by the presentation of the five faces that had also been
353 shown during the encoding and cued recall condition.
354 The faces were shown with two occupations and the
355 correct one had to be selected by button press. One
356 block lasted 21.25 s and included one condition. The
357 given answer was indicated by the font color switch-
358 ing from white to grey. The interstimulus-interval was
359 0.5 s and stimuli were displayed for 3.75 s, regardless
360 of the answer from the participant. After the third run
361 the 15 face-occupation associations were presented
362 again in the same order. This repetition was included
363 to limit the number of stimuli to remember. As stimuli
364 eight pictures of female and seven pictures of male
365 faces from the Ebner face database [48] were shown.

366 As performance measure, we computed task accu-
367 racy for cued recall and recognition blocks.

Spatial construction task

368 Spatial abilities were assessed with a blocked spa-
369 tial construction task (duration: 604 s, Fig. 2). This
370 task was designed in accordance with a task which
371 has been shown to elicit bilateral activation in parietal
372 and occipital brain regions [49]. Spatial abilities are
373 another domain early affected by behavioral deficits
374 in the course of AD [50].

375 The task included four conditions (translation
376 and rotation and the respective control conditions,
377 luminance translation and luminance rotation). Par-
378 ticipants had to either translate or rotate geometric
379 puzzle pieces mentally in order to decide if the pieces
380 fit together to build a square. As control condition,
381 participants indicated if the two grey squares in black
382 boxes in similarly shaped puzzle pieces were of the
383 same grayscale. The task included two runs with eight
384 blocks each. Before each block an instruction text
385 was displayed for 3 s (rotation/translation conditions:
386 “Do these shapes build a square? 2 = Yes, 3 = No”;
387 luminance conditions: “Are the squares displayed
388 in the same opacity? 2 = Same, 3 = Different”). One
389 block lasted 24 s and included one condition. Dur-
390 ing the block the text “2 = Yes, 3 = No” or “2 = Same,
391 3 = Different” was displayed below the puzzle pieces.
392 Each block was presented four times in a fixed
393 order ensuring that translation and rotation conditions
394 would always alternate. Between two blocks, breaks
395 of 12 s and of 27 s between two runs were included
396 during which a black fixation cross was displayed.
397 In the long break (27 s), a text was displayed that
398 now is a short break and the participant should not
399 move. During the task each stimulus was presented
400 for 2.5 s if no button press occurred. After button
401 press or 2.5 s, the stimuli disappeared and a fixation
402 cross appeared for 0.4 s. Stimuli within a block were
403 displayed in a randomized order. Due to the fixed
404 block duration in combination with self-paced trial
405 solving, the number of solved trials differed between
406 participants.

407 Performance was computed analogous to the
408 episodic memory task but here the translation and
409 rotation condition were combined to obtain one value
410 for spatial abilities task accuracy. Due to the fixed
411 block length, the presentation duration of the last trial
412 of each block varied. Therefore, the last trial of each
413 block was not included in the analysis.

414 Participants answered with the index
415 (2/yes/ female/same) and middle finger (3/no/
416 male/different) of the right hand, using a Celeritas®
417 button box with two buttons. Left-handers were
418 asked to use the right-hand.

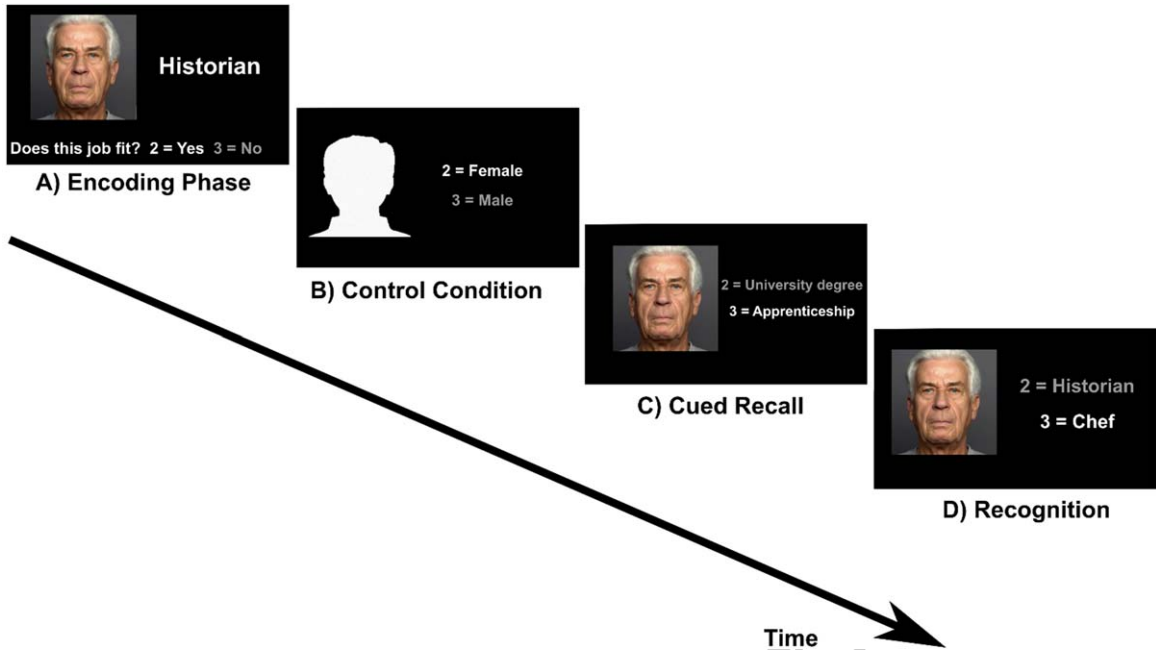


Fig. 1. Face-occupation task. A) Encoding: participants were asked whether a face matches an occupation (subjective rating) B) Control condition: participants were asked to indicate whether a male or female silhouette is shown. C) Cued recall: a face from the encoding condition was presented as cue and participants indicated whether the occupation of the presented person requires a university degree or an apprenticeship. D) Recognition: participants had to choose the correct job between two different options.

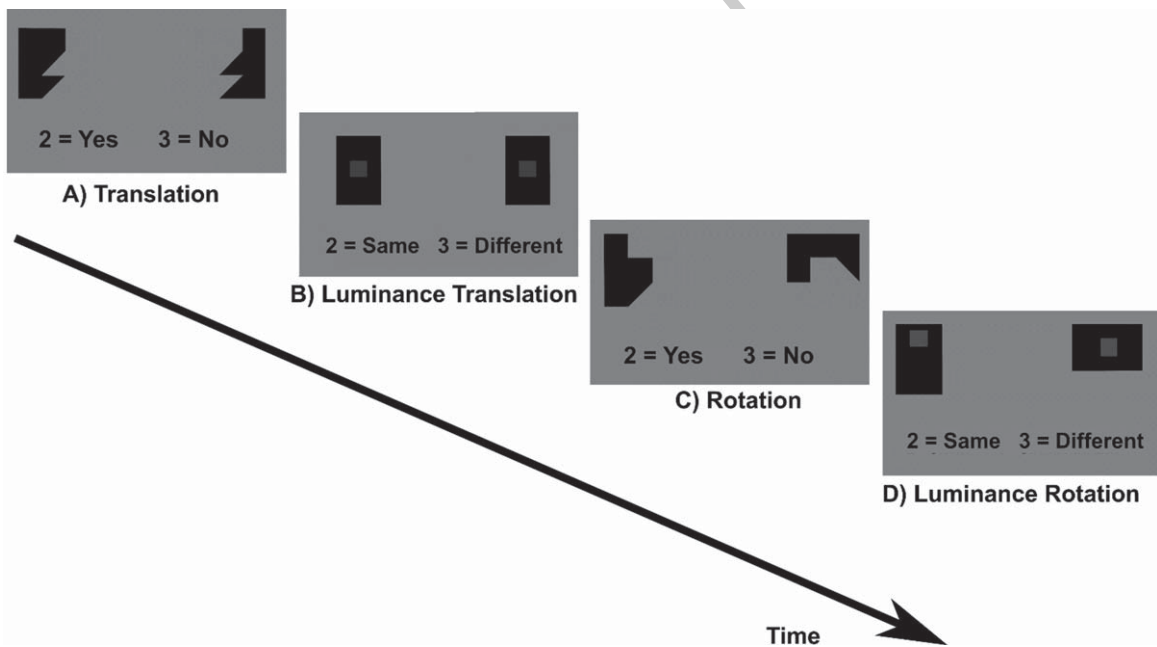


Fig. 2. Spatial construction task. A) Translation and C) Rotation conditions: participants had to either translate or rotate puzzle pieces mentally in order to decide if the pieces fit together to build a square. As control condition, participants indicated if the two grey squares in black boxes in similarly shaped puzzle pieces were of the same grayscale (B) Luminance translation and D) Luminance rotation condition).

419 *MRI data acquisition*

420 Neuroimaging was acquired with a 3T Siemens
 421 scanner (Bern: Siemens Magnetom Prisma, 32 chan-
 422 nel head coil; Lucerne: Siemens Magnetom Vida, 64
 423 channel head coil). Sequences and coil system at the
 424 Lucerne site were adapted to resemble the protocol in
 425 Bern to acquire high quality data while minimizing
 426 hardware differences (please see statistical analy-
 427 ses section for the handling of site differences). The
 428 described MRI parameters below were identical at
 429 both study sites.

430 From all participants, T1- weighted images
 431 (MP2RAGE, TR = 5000 ms, TE = 2.98 ms, TI
 432 1/2 = 700 ms/2500 ms, flip angle 1/2 = 4°/5°,
 433 FOV = 256 mm x 256 mm, matrix = 256 x 256, voxel
 434 size = 1 x 1 x 1 mm, 176 slices) were acquired.

435 For the fMRI sequences we used echo-planar
 436 imaging (EPI) with 604 (spatial abilities Task) and
 437 638 (episodic memory task) volumes (TR = 1000 ms,
 438 TE = 37 ms, flip angle = 30°, FOV = 230 mm x
 439 230 mm, matrix = 92 x 92, accelerating factor 8, voxel
 440 size 2.5 x 2.5 x 2.5 mm, 72 slices). The axial slices
 441 were positioned along the anterior commissure and
 442 the posterior commissure.

443 *MRI processing*

444 To calculate hippocampal atrophy, the struc-
 445 tural images were automatically segmented with
 446 the computational anatomy toolbox (CAT12:
 447 <http://www.neuro.uni-jena.de/cat/>). ROIs were
 448 calculated using the neuromorphometrics atlas. To
 449 calculate hippocampal atrophy, total hippocampal
 450 volume was scaled by total intracranial volume
 451 and the results subtracted from one, so that larger
 452 numbers were corresponding to higher levels of
 453 hippocampal atrophy.

454 To provide a grey matter mask for the task fMRI
 455 group analyses, the anatomical images of each partic-
 456 ipant were segmented using SPM12 [51] in MATLAB
 457 version 2019b (Natick, MA: The MathWorks Inc.) on
 458 a Linux platform. After segmentation the individual
 459 grey matter images were spatially normalized to stan-
 460 dard MNI space and then combined into one mean
 461 image over all participants as grey matter mask.

462 Functional volumes were first realigned to the
 463 mean image of each individual, coregistered to
 464 the anatomical image in native space and finally
 465 smoothed with a 6 mm full width at half maximum
 466 Gaussian Kernel.

467 As first-level analyses general linear models were
 468 computed for each participant in native space with
 469 BOLD signal changes as dependent variable. Each
 470 block type was included as separate predictors with
 471 one additional predictor for instruction screens and
 472 breaks (episodic memory task: encoding, control,
 473 cued recall, recognition, instruction screens; spatial
 474 abilities task: translation, rotation, luminance trans-
 475 lation, luminance rotation, instruction screens). To
 476 account for possible head movements, the absolute
 477 values of the first derivate of the six default move-
 478 ment parameters obtained during realignment were
 479 included. Each predictor's time course was convolved
 480 with a standard hemodynamic response function.
 481 A 128-s high pass filter was used to account for
 482 scanner drift and a separate variable was added to
 483 model the intercept. The resulting first level con-
 484 trast images were then normalized into MNI space
 485 and resampled to isometric voxels with a side length
 486 of 2 mm.

487 To test if severe motion artefacts were present
 488 in the fMRI data, the absolute derivate of the
 489 first three movement parameters (x, y, z axis)
 490 were checked for values exceeding 2 mm between
 491 subsequent volumes. Two participants exceeded
 492 this threshold in the episodic memory task, one
 493 participant 17 times and one participant once.
 494 The participant with 17 movements was excluded
 495 from data analysis in the face-occupation episodic
 496 memory task. In the spatial abilities task, move-
 497 ments exceeding 2 mm were observed once in
 498 one participant and, therefore, no participants were
 499 excluded.

500 *Statistical analyses*

501 *Behavioral data analyses*

502 To investigate the association between blood-based
 503 biomarkers, behavioral composite scores and hip-
 504 pocampal atrophy we performed correlation analyses
 505 as well as partial Pearson's correlation analyses con-
 506 trolling for age. For categorical variables (APOE ϵ 4
 507 carrier: yes/no) Welch's t-tests were used. We also
 508 used Welch's t-tests to test for significant differences
 509 between study sites in behavioral composite scores,
 510 hippocampal atrophy, and fMRI task accuracy. Paired
 511 t-tests were used to investigate differences in task
 512 accuracy, reaction times and number of solved trials
 513 between the different conditions in the fMRI tasks.

514 The analysis of the behavioral data was performed
 515 with R studio [52].

Imaging data analyses

Spatially normalized contrast images from the individual subjects level coding cued recall > control contrasts (i.e., deciding if the occupation associated with the presented face requires an apprenticeship or university degree>is the presented head contour male or female) from the face-occupation episodic memory task entered group level analyses (second level). We expected this contrast to elicit parietal activity associated with episodic memory retrieval processes. A study using a similar face-name paradigm for example, reported activation during successful retrieval in the posteromedial cortex compared to lower activation during encoding in young adults [53]. Furthermore, cued recall tasks are sensitive markers for early AD related memory impairment [13, 47]

For the visual construction task, we selected the [translation+rotation]>[luminance translation+luminance rotation] contrast, in accordance with the results of Seydell et al. [49] which showed strong parietal activity.

As outlined in the introduction, patterns reflecting neuronal compensation were defined as regions where greater functional activity is associated with greater hippocampal atrophy and better task performance [27].

To investigate potential neuronal compensation, we used inclusive masking with an orthogonal (i.e., independent) contrast as in previous work [54]. In detail, significant activity with an uncorrected threshold of $p < 0.01$ in a first t-contrast for activity correlating with hippocampal atrophy was calculated. The resulting image was used as inclusive binary mask for a second estimation of the same contrast with activity positively correlated with fMRI task performance. For this second contrast estimation a family-wise error correction (FWE, $p < 0.05$) for multiple testing as well as a less conservative threshold of $p < 0.001$ uncorrected were used. The masking image included only few clusters where activity could be detected in the second contrast. Therefore, p-values in the second contrast estimation were corrected for the small search region with a small volume correction after masking.

While we selected tasks especially activating parietal areas, compensatory brain activity might also appear in brain areas outside the parietal lobe. Therefore, we performed whole brain analyses with the grey matter mask calculated for the study sample (please see below for detailed information about the grey matter mask). We included the fMRI in-task per-

formance measures in the main models (model 1.1: Compensation related to performance in the episodic fMRI task; model 2.1: Compensation related to performance in the spatial abilities fMRI task). In a secondary analysis, we included performance measures from the behavioral composite scores instead of fMRI task performance, these models are reported in the Supplementary Material (model 1.2: Compensation related to the memory composite score; model 2.2: Compensation related to the spatial abilities composite score).

As previously mentioned, we repeated all analyses with a subsample with positive blood biomarkers for amyloid positivity (model 1.3: Compensation related to performance in the episodic memory fMRI task in a subsample with positive blood biomarkers for amyloid positivity; model 2.3: Compensation related to performance in the spatial abilities fMRI task in a subsample with positive blood biomarkers for amyloid positivity). The results from the subsample whole brain analyses and the detailed results from the models including fMRI task performance are reported in the Supplementary Material (model 1.4: Compensation related to performance in the memory composite score in a subsample with positive blood biomarkers for amyloid positivity; model 2.4: Compensation related to the spatial abilities composite score in a subsample with positive blood biomarkers for amyloid positivity). We did not correct for the number of models as we aimed to detect indications for compensation and consequently employed liberal statistical thresholds throughout (i.e., $p < 0.01$ for the masking image and $pFWE < 0.05$ as well as $p_{\text{uncorrected}} < 0.001$ for the combination of both contrast images) [55]. For the same reason, we did not apply a voxel extent threshold. In all models one-sample t-tests were performed. Another possibility to investigate neuronal compensation is to build two samples according to the residuals in a linear regression for task performance and hippocampal atrophy. Participants with positive residuals in the regression scored better than estimated based on their hippocampal volume and we expect neuronal compensation to be most likely in this group (i.e., sample one). This is not the case in participants with null or negative residuals (i.e., sample two). We compared these samples with two-sample t-tests. Furthermore, we performed two-sample t-tests to compare subsamples with and without probable amyloid positivity based on blood biomarkers. The results for these analyses are reported in the Supplementary Material.

Table 1
Demographics of participants

	Mean	SD	Range
Age (y)	71	5.70	60–81
Gender (m/f)	22/30		
Education (y)	15.15	3.06	9–20
MoCA Score	27.32	1.97	24–30
Composite Score M	0.00	0.93	–2.29–1.6
Composite Score SA	0.00	0.49	–1.20–0.93

SD, standard deviation; M, memory; SA, spatial abilities.

All reported p -values in the fMRI analyses correspond to the peak-level significance and coordinates to the MNI space (x, y, z).

Age was related to hippocampal atrophy and behavioral performance (please see the section “Relationship between behavioral data and hippocampal atrophy” for the results of the correlation analyses) and, therefore, was included as covariate in all fMRI analyses. Because MRI devices and head coils were different between study sites, site was also included as covariate. Sex was not related to hippocampal atrophy or behavioral composite scores (please see Table 3 for detailed results), but to address potential neuronal differences between females and males [56], sex was included as additional covariate.

RESULTS

Demographic details for the participants are summarized in Table 1. The mean score of the geriatric depression scale [42] was 1.73 (SD: 1.55). No participant had ten or more points, which would indicate severe depressive symptoms.

The mean values and standard deviations of the six test scores included in the behavioral composite scores are reported in the Supplementary Material.

To explore potential effects of study site on behavioral outcomes and hippocampal atrophy scores Welch’s t -tests were performed for the behavioral composite scores (memory: $t(35.68) = -1.97, p = 0.06$; spatial abilities: $t(31.95) = -0.1, p = 0.91$), hippocampal atrophy ($t(46.59), p = 0.50$) and fMRI task accuracy (memory: $t(23.71), p = 0.72$; spatial abilities: $t(32.13), p = 0.93$). The results did not indicate significant differences between study sites in any of these outcomes. Therefore, the correlation analyses were not controlled for study site.

Composite scores

For the episodic memory outcomes, the PCA showed very similar loadings (loading AVLT delayed

recall: -0.58 , loading immediate recall: -0.57 , loading total learning sum -0.58). Therefore, no weighting of the single raw test scores was performed for the behavioral composite score. The PCA for the SA raw scores showed differences between the loadings of the encoding and the two other scores which were larger than 0.05 (loading RCF encoding: -0.41 , loading immediate recall: -0.65 , loading delayed recall: -0.6). Therefore, we weighted the individual test scores with their absolute loads before combining them into one composite score.

Relationship between behavioral data and hippocampal atrophy

There was a significant correlation between performance in measures of episodic memory and hippocampal atrophy (fMRI task accuracy: $r = -0.32, p < 0.05$; memory composite score: $r = -0.34, p = 0.01$) indicating that higher performance in the episodic memory task or memory composite score was associated with lower hippocampal atrophy (Fig. 3). There were no correlations between measures of spatial abilities tasks and hippocampal atrophy (fMRI task performance: $r = -0.07, p = 0.62$; spatial abilities composite score: $r = -0.25, p = 0.07$).

Age was significantly correlated with hippocampal atrophy ($r = 0.46, p < 0.001$), episodic memory fMRI task performance ($r = -0.49, p < 0.001$) and the composite scores (memory: $r = -0.31, p < 0.05$; spatial abilities: $r = -0.29, p < 0.05$). This indicated higher hippocampal atrophy and lower task performance with increasing age. There was no significant correlation between performance in the spatial abilities fMRI task and age ($r = -0.21, p = 0.14$).

Cook’s distance plots did not indicate any influential data points (Cook’s distances larger than 0.5) in the correlations.

When controlled for age, no correlations were significant (episodic memory task performance and hippocampal atrophy: $r = -0.12, p = 0.39$; memory composite score and hippocampal atrophy: $r = -0.24, p = 0.09$).

Blood-based biomarkers

Seven participants were carriers of at least one APOE $\epsilon 4$ allele. Furthermore, we analyzed A $\beta_{42/40}$ ratio (mean = 0.068, SD = 0.014), pTau181 (mean = 2.34 pg/ml, SD = 1.22), glial fibrillary acidic protein (mean = 126.86 pg/ml, SD = 60.21),

Table 3

Association between blood-based biomarkers, behavioral composite scores, sex, and hippocampal atrophy (n=38). For the dichotomous variable APOE $\epsilon 4$ carrier (yes/no) and sex (female/male) Welch's t-tests were performed and t-values are reported. For the other blood-based biomarkers partial Pearson's correlations were calculated and partial correlation coefficients are reported

Blood-based biomarkers	Memory composite score	Spatial abilities composite score	Hippocampal atrophy
APOE4 carrier (Welch's t)	0.00	1.14	0.24
Sex (Welch's t)	-1.2	1.37	0.77
Amyloid- $\beta_{42/40}$ ratio (partial r)	0.20	0.06	-0.55**
pTau181 (partial r)	0.07	-0.13	-0.13

* $p < 0.05$, ** $p < 0.001$.

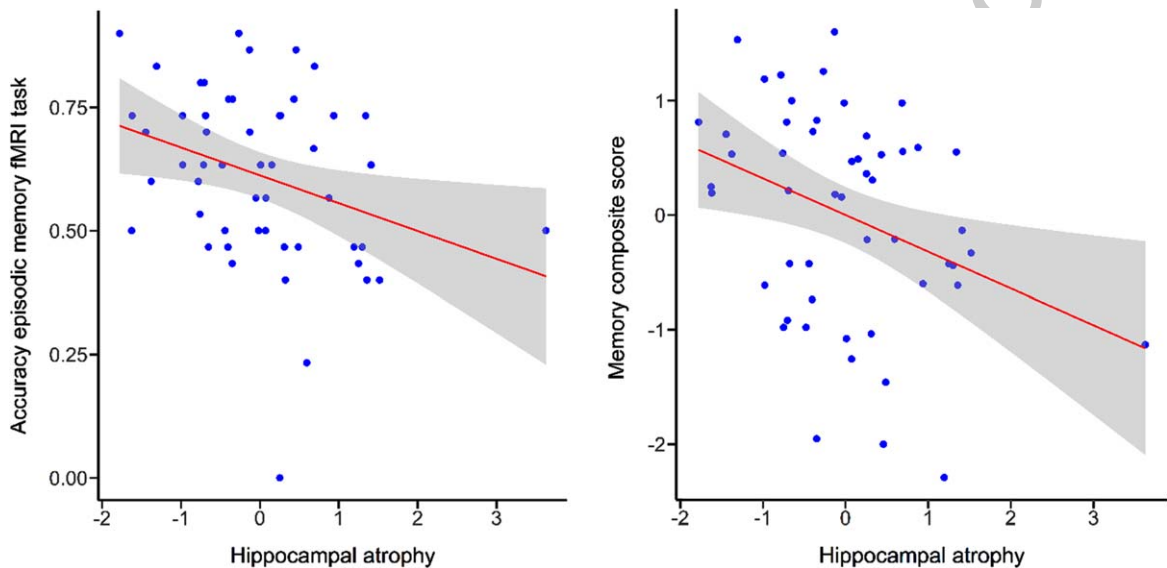


Fig. 3. Scatterplots illustrating correlations for episodic memory fMRI task performance and memory composite score with hippocampal atrophy. Grey bands indicate standard errors.

706 and neurofilament light chain (mean = 22.9 pg/ml, SD = 11.21). Based on currently recommended cut-off scores, ratios lower than 0.06 in the $A\beta_{42/40}$ ratio indicate amyloid positivity [57]. In our sample, this included eight participants. For pTau181, a cut-off of 1.8 pg/ml has been suggested for amyloid positivity [57], which indicated 18 participants with amyloid positivity. This pTau181 cut-off score is in-line with previous research [58]. Three participants reached the cut-off scores for amyloid positivity in both the $A\beta_{42/40}$ ratio and pTau181. Therefore, we considered 23 participants as positive for AD pathology related blood-based biomarkers. There was a significant association between hippocampal atrophy and $A\beta_{42/40}$ ratio ($r = -0.55$, $p < 0.001$) when controlling for age. Please see Table 3 for the complete results of the performed Welch's t-test and partial Pearson's correlation analyses.

724 The correlation between $A\beta_{42/40}$ ratio and hippocampal atrophy remained significant also when the correlation analysis was additionally controlled for sex and years of education ($r = -0.60$, $p < 0.001$).

728 *fMRI task performance*

729 The accuracy levels across all conditions in the two fMRI tasks were significantly above the chance level of 50% (cued recall: $t(51) = -4.59$, $p < 0.001$; recognition: $t(51) = -8.71$, $p < 0.001$, translation: $t(51) = -10.58$, $p < 0.001$; rotation: $t(51) = -10.40$, $p < 0.001$; luminance translation: $t(51) = -11.49$, $p < 0.001$; luminance rotation: $t(51) = -10.54$, $p < 0.001$). Therefore, we assume that both tasks were appropriately designed regarding task duration and level of difficulty.

Model 1: Episodic memory fMRI task

A whole brain analysis (constrained to the grey matter mask) for the cued recall > control contrast in the face-occupation episodic memory fMRI task showed significant activity in the parietal and occipital lobe. Therefore, we considered the episodic memory task and the selected contrast as appropriate and conducted further analyses (please see the Supplementary Material for details). Study site, total intracranial volume, age, and sex were included as covariates.

Model 1.1: Compensation related to performance in the episodic memory fMRI task

A mask for task-related activity positively correlated with hippocampal atrophy ($p_{\text{uncorrected}} < 0.01$) was calculated in a first step in the cued recall > control contrast (Fig. 4A).

In a second step, the same contrast was calculated, but this time for activity that was positively correlated with fMRI task performance (i.e., accuracy). Task accuracy was not correlated with significant brain activity in any brain region when corrected for multiple testing ($p_{\text{FWE}} < 0.05$). Without correction for multiple testing, the strongest activity before masking was located in the left lateral orbital gyrus ($t = 3.49$; $p_{\text{uncorrected}} = 0.001$; peak x, y, z coordinates: -40, 40, -18) and the left cerebral white matter/occipital fusiform gyrus (Fig. 4B).

Finally, the binary mask was used (inclusive masking) in the same contrast for activity that was positively correlated with episodic memory fMRI task performance. There were no significant clusters after masking ($p_{\text{FWE}} < 0.05$ or $p_{\text{uncorrected}} < 0.001$).

Model 1.2: Compensation related to the memory composite score

An identical inclusive masking analysis with the memory composite score instead of episodic memory fMRI task accuracy did not show any significant results ($p_{\text{FWE}} < 0.05$ or $p_{\text{uncorrected}} < 0.001$). Please see the Supplementary Material for details.

Model 2: Spatial abilities fMRI task

A whole brain analysis ([translation+rotation]>luminance conditions contrast, constrained to the grey matter mask) with study site, total intracranial volume and age as covariates was calculated and revealed several clusters with significant activity in the parietal lobe (please see

the Supplementary Material for details). Therefore, we considered the selected spatial abilities contrast as appropriate for the planned analyses.

Model 2.1: Compensation related to performance in the spatial abilities fMRI task

We calculated also for the spatial abilities fMRI paradigm task-related activity positively correlated with hippocampal atrophy ($p_{\text{uncorrected}} < 0.01$). This showed the strongest activity in the left subcallosal area ($t = 4.78$; $p_{\text{uncorrected}} = 0.000$; peak x, y, z coordinates: -4, 10, -24), cerebellar vermal lobules I-V ($t = 4.18$; $p_{\text{uncorrected}} = 0.000$; peak x, y, z coordinates: 0, -54, -16), and the left cerebral white matter/superior frontal gyrus ($t = 3.91$; $p_{\text{uncorrected}} = 0.000$; peak x, y, z coordinates: -20, 12, 46) (Fig. 5A).

Task accuracy was not positively correlated with significant brain activity when corrected for multiple testing ($p_{\text{FWE}} < 0.05$). Without correction for multiple testing ($p_{\text{uncorrected}} < 0.001$) the only significant clusters were located in the right medial orbital gyrus ($t = 4.62$; $p_{\text{uncorrected}} = 0.000$; peak x, y, z coordinates: 14, 26, -30; $t = 3.91$; $p_{\text{uncorrected}} = 0.000$; peak x, y, z coordinates: 18, 18, -28; $t = 3.78$; $p_{\text{uncorrected}} = 0.000$; peak x, y, z coordinates: 12, 20, -28) and the left supra-marginal gyrus ($t = 3.59$; $p_{\text{uncorrected}} = 0.000$; peak x, y, z coordinates: -66, -32, 32) (Fig. 5B).

After inclusive masking, one isolated effect with significant activity was located in the left supra-marginal gyrus without correction for multiple testing ($t = 3.64$; $p_{\text{uncorrected}} = 0.000$; peak x, y, z coordinates: -66, -32, 32) (Fig. 5C).

Model 2.2: Compensation related to the spatial abilities composite score

A model with hippocampal atrophy as mask ($p_{\text{uncorrected}} < 0.01$) for activity positively correlated with the spatial abilities composite score also did not show any significant results ($p_{\text{FWE}} < 0.05$ or $p_{\text{uncorrected}} < 0.001$). Please see the Supplementary Material for details.

Repetition of fMRI analyses in a subsample with positive blood biomarkers for amyloid positivity

Model 1.3: Compensation related to performance in the episodic memory fMRI task in a subsample with positive blood biomarkers for amyloid positivity

A contrast for the positive correlation between activity and hippocampal atrophy was calculated

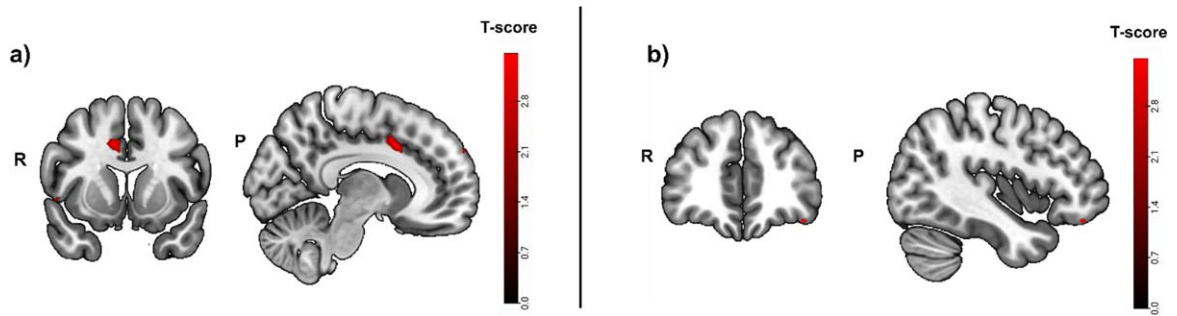


Fig. 4A. Greater activity related to greater hippocampal atrophy in the episodic memory task (cued recall>control contrast, $p_{uncorrected} < 0.01$), used as mask. Activity was detected in the left temporal pole and the right middle cingulate and occipital fusiform gyrus. b) Greater activity related to high episodic memory task accuracy in the cued recall>control contrast was located in the left lateral orbital gyrus and the left cerebral white matter/occipital fusiform gyrus ($p_{uncorrected} < 0.001$, before masking). R, right; P, posterior.

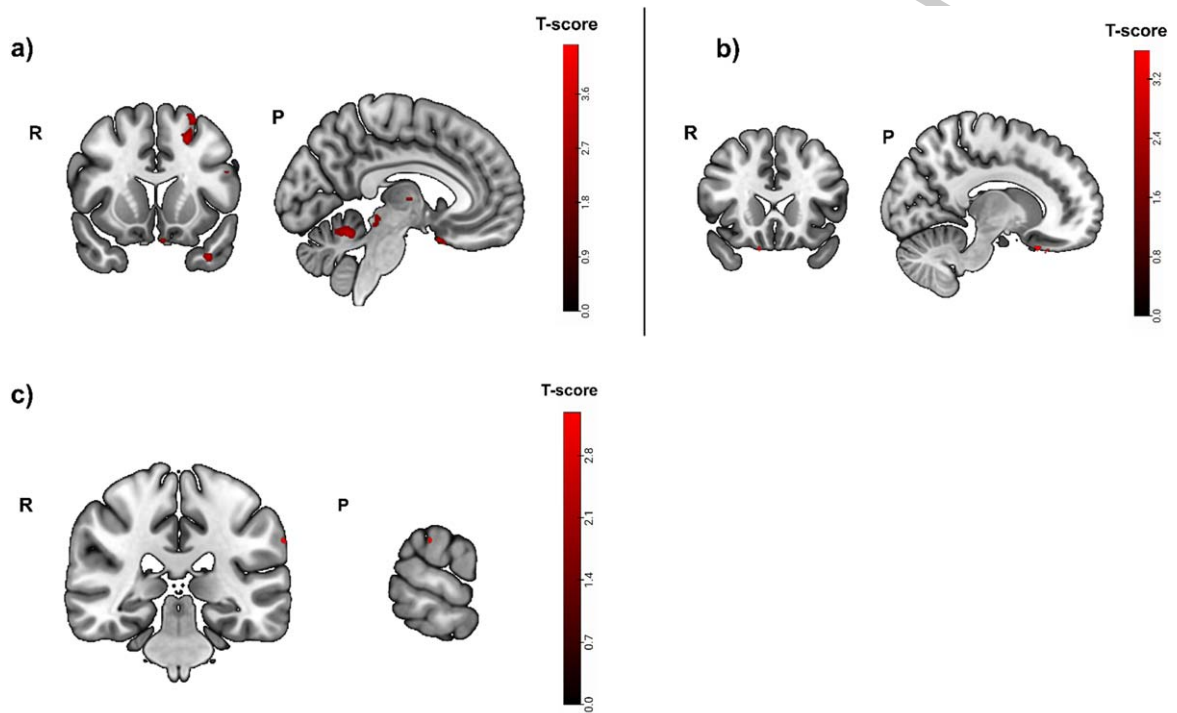


Fig. 5A. Greater activity related to greater hippocampal atrophy in the spatial abilities task ([translation+rotation]>luminance conditions contrast, $p_{uncorrected} < 0.01$), used as mask. Activity was detected in the left subcallosal area, the cerebellar vermal lobules I-V and the left cerebral white matter/superior frontal gyrus. b) Greater activity related to high spatial abilities task accuracy (before masking) was located in the right medial orbital gyrus and the left supramarginal gyrus ($p_{uncorrected} < 0.001$). c) After inclusive masking a significant cluster in the left supramarginal gyrus was detected ($p_{uncorrected} < 0.001$). R, right; P, posterior.

834 ($p_{uncorrected} < 0.01$) (Fig. 6A). The clusters with
 835 strongest activity were detected in the left middle
 836 temporal gyrus ($t = 5.50$; $p_{uncorrected} = 0.000$; peak x,
 837 y, z coordinates: -66, -10, -20), the right angular gyrus
 838 ($t = 5.41$; $p_{uncorrected} = 0.000$; peak x, y, z coordinates:
 839 48, -64, 18) and left cerebral white matter/temporal
 840 pole ($t = 5.27$; $p_{uncorrected} = 0.000$; peak x, y, z coord-
 841 inates: -44, 4, -22).

842 Task accuracy was not positively correlated to sig-
 843 nificant brain activity when corrected for multiple
 844 testing ($p_{FWE} < 0.05$). Without correction for mul-
 845 tiple testing, the strongest activity was located in the
 846 right calcarine cortex ($t = 4.80$; $p_{uncorrected} = 0.000$;
 847 peak x, y, z coordinates: 8, -84, -2), the left middle
 848 frontal gyrus ($t = 4.70$; $p_{uncorrected} = 0.000$; peak x, y,
 849 z coordinates: -26, 6, 56) and the left cerebral white

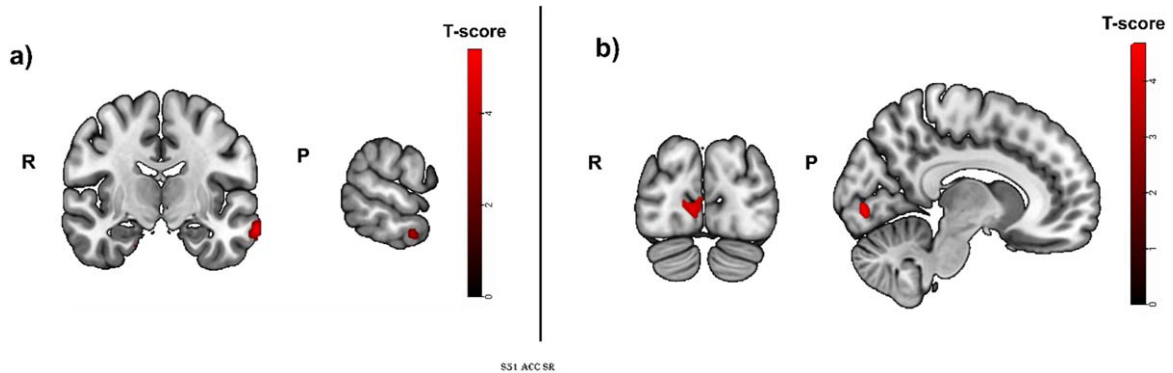


Fig. 6A. Greater activity related to greater hippocampal atrophy in the episodic memory task (cued recall > control contrast, $p_{\text{uncorrected}} < 0.01$), used as mask in a subsample with positive blood biomarkers for amyloid positivity. Activity was detected in the left middle temporal gyrus, the right angular gyrus and left cerebral white matter/temporal pole. b) Greater activity related to high episodic memory task accuracy (cued recall > control contrast, before masking) was located in the right calcarine cortex, the left middle frontal gyrus, the left cerebral white matter and opercular part of the inferior frontal gyrus. R, right; P, posterior.

matter/opercular part of the inferior frontal gyrus ($t = 4.46$; $p_{\text{uncorrected}} = 0.000$; peak x, y, z coordinates: -48, 16, 16) (Fig. 6B).

Also in this contrast no voxel survived when the binary mask for hippocampal atrophy was applied ($p_{\text{uncorrected}} < 0.001$).

Model 1.4: Compensation related to the memory composite score in a subsample with positive blood biomarkers for amyloid positivity

A model with hippocampal atrophy as mask ($p_{\text{uncorrected}} < 0.01$) for activity positively correlated with episodic memory task accuracy did not show any significant results ($p_{\text{FWE}} < 0.05$ or $p_{\text{uncorrected}} < 0.001$). Please see the Supplementary Material for details.

Model 2.3: Compensation related to performance in the spatial abilities fMRI task in a subsample with positive blood biomarkers for amyloid positivity

The [translation+rotation]>luminance conditions contrast showed the strongest activity ($p_{\text{uncorrected}} < 0.01$) positively correlated with hippocampal atrophy in the left middle frontal gyrus ($t = 5.24$; $p_{\text{uncorrected}} = 0.000$; peak x, y, z coordinates: -40, 2, 60), the left superior frontal gyrus ($t = 4.95$; $p_{\text{uncorrected}} = 0.000$; peak x, y, z coordinates: -24, 10, 66), and the left superior frontal gyrus medial segment ($t = 4.81$; $p_{\text{uncorrected}} = 0.000$; peak x, y, z coordinates: -6, 64, 24) (Fig. 7A).

No significant results remained after $p_{\text{FWE}} < 0.05$ correction when positive correlations with the spatial abilities task accuracy were calculated. Without cor-

rection for multiple testing ($p_{\text{uncorrected}} < 0.001$) the only significant cluster was located in the right medial orbital gyrus ($t = 4.56$; $p_{\text{uncorrected}} = 0.000$; peak x, y, z coordinates: 14, 26, -30) and close by in the gyrus rectus ($t = 3.68$; $p_{\text{uncorrected}} = 0.001$; peak x, y, z coordinates: 6, 24, -32) (Fig. 7B).

After inclusive masking no significant voxel survived ($p_{\text{uncorrected}} < 0.001$).

Model 2.4: Compensation related to the spatial abilities composite score in a subsample with positive blood biomarkers for amyloid positivity

A model with hippocampal atrophy as mask ($p_{\text{uncorrected}} < 0.01$) for activity positively correlated with the spatial abilities composite score also did not show any significant results ($p_{\text{FWE}} < 0.05$ or $p_{\text{uncorrected}} < 0.001$). Please see the Supplementary Material for details.

DISCUSSION

In the present study, we investigated if neuronal compensation existed in a sample of older adults with SCD. We employed two fMRI tasks targeting different cognitive domains. Both tasks should induce activity in the parietal lobe, which is affected by neuropathology early in the course of AD.

We did not find strong evidence for compensatory brain activity in either of the two tasks. The model for the complete sample in the spatial abilities task (model 2.1) showed a very small effect in the left supramarginal gyrus uncorrected for multiple comparisons. This brain region has been associated with working memory and attention. It is part of the fronto-

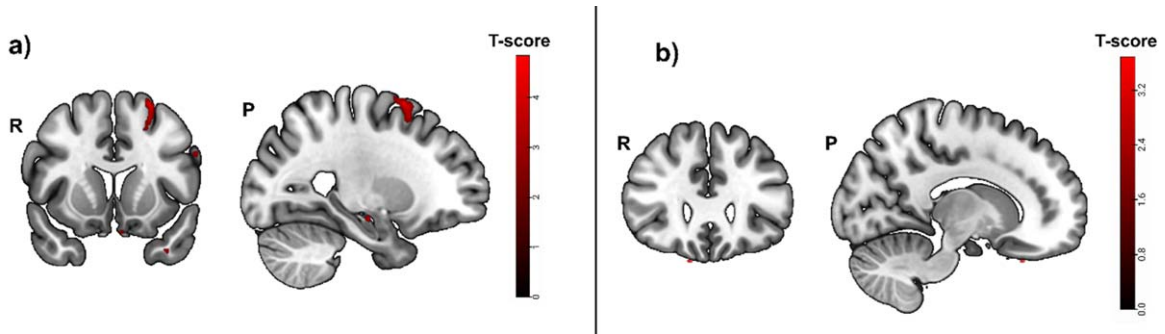


Fig. 7A. Greater activity related to greater hippocampal atrophy in the spatial abilities task ([translation+rotation]>luminance conditions contrast, $p_{uncorrected} < 0.01$), used as mask in a subsample with positive blood biomarkers for amyloid positivity. Activity was detected in the left middle frontal gyrus, the left superior frontal gyrus and the left superior frontal gyrus medial segment. b) Greater activity related to high spatial abilities task accuracy (before masking) was located in the right medial orbital gyrus and the gyrus rectus ($p_{uncorrected} < 0.001$).

parietal attentional control network and crucial for spatial working memory [59]. Another study reported compensatory activity in the left parietal lobe in early neurodegeneration [11], which is located closely to the effect we detected.

However, in fMRI studies, a common minimal cluster size is ten voxels [59]. Regarding the very small size of three voxels, the small t-value ($t = 3.64$) and the location of our finding at the outer grey matter/CSF border, this result must be interpreted with caution. In a subsample with blood biomarkers indicating amyloid positivity, and in subsamples based on residuals of regressions of task performance and hippocampal atrophy, we found no evidence for neuronal compensation.

Whole brain analysis for the contrast cued recall>control in the face-occupation task showed the strongest activity in the occipital lobe and the left ventral diencephalon. The activity over the occipital lobe was partially caused by the visually different stimuli in both conditions [46]. The cluster including the left diencephalon encompassed also structures like the parahippocampal gyrus and the hippocampus, which can be expected when using a task targeting episodic memory [16]. The whole brain analysis for the [translation+rotation]>luminance conditions contrast in the spatial abilities task showed the strongest activation in the inferior occipital gyrus encompassing several structures from the parietal and occipital lobe in both hemispheres. This corresponds to the activity pattern reported for young adults in this task [49]. Our results show that the selection of tasks and contrasts was appropriate for eliciting activity in the parietal cortex.

Previous research supports the existence of neuronal compensation [11, 13, 14, 26]. However, there

are substantial differences between these and the present study, which might explain the different findings. The other studies included different samples for healthy controls and participants affected by SCD [11, 13, 14] and MCI [26]. To fulfil the criteria of successful compensation, greater activity has to be related to better performance. In studies including SCD as well as healthy controls this correlation would appear in the SCD sample only [60]. This has been the case in the studies from Erk et al. [13] and Corriveau-Lecavalier et al. [11, 61]. As only these two studies fulfil the criteria for successful compensation in SCD our results indicate that compensation in this population is not a stable finding.

While the absence of clear compensatory brain activity in our full sample might be due to the inclusion of subjects without neuropathology this explanation is less likely in the subset with positive blood biomarkers for amyloid positivity. For the identification of amyloid positivity, we relied on relatively new blood biomarkers for amyloid- β and pTau181. The $A\beta_{42/40}$ ratio has shown to be a good measure for amyloid PET status [62]. In our sample, eight participants were positive for amyloid pathology according to the cut-off score. To increase sample size, we also included participants with high plasma pTau181 values. This blood biomarker predicted amyloid PET positivity [63]. It is possible that current plasma pTau181 measures are not sensitive enough to differentiate within a sample of SCD for amyloid positivity yet. This assumption is supported by the lack of a correlation between pTau181 values and our measure of hippocampal atrophy ($r = -0.13$, $p = 0.44$), which was correlated with the $A\beta_{42/40}$ ratio ($r = -0.55$, $p < 0.001$) also when corrected for age.

985 It is possible that different patterns of brain activity
 986 in neuronal compensation occur in our study sample.
 987 Therefore, it might be interesting to investigate single
 988 subject data to identify groups with different patterns
 989 of compensatory activity in a future study. Another
 990 statistical approach would be to investigate fMRI data
 991 with dimensional reduction methods following, e.g.,
 992 an independent component analysis [64]. It is pos-
 993 sible that such alternative approaches would lead to
 994 different results but with the downside of inflating the
 995 number of tests.

996 *Limitations*

997 We included no sample of healthy subjects, which
 998 might have facilitated the detection of neuronal com-
 999 pensation.

1000 We used hippocampal volume as a proxy for dis-
 1001 ease load and therefore refer to it as hippocampal
 1002 atrophy. This term reflects our interpretation of vol-
 1003 ume, but we have no knowledge on longitudinal
 1004 changes in hippocampal size of participants. There-
 1005 fore, it is possible that some participants with high
 1006 hippocampal atrophy according to our study results
 1007 were actually born with a relatively small hippocam-
 1008 pus and no atrophy occurred. Data for blood-based
 1009 biomarkers was available for 38 participants, result-
 1010 ing in a final sample of 23 participants with biomarker
 1011 values indicating amyloid positivity. This sample size
 1012 might have been too small to find subtle effects of
 1013 early AD related pathology in brain activity. Addi-
 1014 tionally, in our sample only eight participants were
 1015 positive for amyloid pathology according to the cut-
 1016 off score for the $A\beta_{42/40}$ ratio. A larger number might
 1017 have been necessary to detect neuronal compensa-
 1018 tion. This limitation can also be applied to the sample
 1019 as a whole.

1020 As mentioned in the introduction, SCD is non-
 1021 specific and can appear as an early sign of cognitive
 1022 decline but also as consequence of psychiatric dis-
 1023 orders [2]. Therefore, our sample probably included
 1024 SCD due to different causes, while compensatory
 1025 activity in SCD is mostly expected to be a result
 1026 of beginning neurodegeneration. This heterogeneity
 1027 might have reduced our ability to detect strong com-
 1028 pensatory brain activity on the group level.

1029 While our approach focused on increased brain
 1030 activity, also reduced activity and network connec-
 1031 tivity have been observed in SCD [65]. The authors
 1032 of this review suggest a model where neuronal con-
 1033 nectivity is related to SCD stage (i.e., after increased
 1034 connectivity due to noisy signal propagation and

potential compensation in early SCD connectivity
 decreases). In our sample we considered SCD as one
 category and did not include the onset of SCD as
 potential moderating factor of compensatory effects.
 But we expect hippocampal atrophy as an indicator
 of disease load to increase in the course of SCD.

1041 *Conclusion*

1042 Our study did not provide conclusive evidence
 1043 for compensatory brain activity in older adults with
 1044 SCD in tasks targeting episodic memory or spatial
 1045 abilities. It is possible that SCD is too early in the
 1046 process of neurodegeneration to elicit compensatory
 1047 activity or that this activity is too divergent among
 1048 individuals given the broad definition of SCD in com-
 1049 bination with the sample size used. Future studies
 1050 could emphasize on detecting compensation in the
 1051 individual as interventions such a transcranial electric
 1052 current stimulation could be adapted to the individ-
 1053 ual's pattern of compensation.

1054 **ACKNOWLEDGMENTS**

1055 We would like to thank Tania Geiger, Dominique
 1056 Hürzeler, Julia Hübscher, Fiona Pfister, and Raphaela
 1057 Zwimpfer for their help with the cognitive assess-
 1058 ments. Additionally, we would like to thank all
 1059 participants for their commitment.

1060 **FUNDING**

1061 This research was funded by the Swiss
 1062 National Science Foundation (SNSF; grant number
 1063 32003B_189240).

1064 **CONFLICT OF INTEREST**

1065 The authors declare that the research was con-
 1066 ducted in the absence of any commercial or financial
 1067 relationships that could be construed as a potential
 1068 conflict of interest.

1069 **DATA AVAILABILITY**

1070 The relevant data for this publication is openly
 1071 available in the "Bern Open Repository and Infor-
 1072 mation System" at <https://doi.org/10.48620/66>.

SUPPLEMENTARY MATERIAL

The supplementary material is available in the electronic version of this article: <https://dx.doi.org/10.3233/JAD-221001>.

REFERENCES

- [1] Zullo L, Clark C, Gholam F, Castelao E, von Gunten A, Preisig M, Popp J (2021) Factors associated with subjective cognitive decline in dementia-free older adults-A population-based study. *Int J Geriatr Psychiatry* **36**, 1188-1196.
- [2] Jessen F, Amariglio RE, Buckley RF, van der Flier WM, Han Y, Molinuevo JL, Rabin L, Rentz DM, Rodriguez-Gomez O, Saykin AJ, Sikkes SAM, Smart CM, Wolfsgruber S, Wagner M (2020) The characterisation of subjective cognitive decline. *Lancet Neurol* **19**, 271-278.
- [3] Hao L, Xing Y, Li X, Mu B, Zhao W, Wang G, Wang T, Jia J, Han Y (2019) Risk factors and neuropsychological assessments of subjective cognitive decline (plus) in Chinese memory clinic. *Front Neurosci* **13**, 846.
- [4] Koppara A, Wagner M, Lange C, Ernst A, Wiese B, König H-H, Brettschneider C, Riedel-Heller S, Lupp A, Weyerer S, Werle J, Bickel H, Mösch E, Pentzek M, Fuchs A, Wolfsgruber S, Beauducel A, Scherer M, Maier W, Jessen F (2015) Cognitive performance before and after the onset of subjective cognitive decline in old age. *Alzheimers Dement (Amst)* **1**, 194-205.
- [5] Pike KE, Cavuoto MG, Li L, Wright BJ, Kinsella GJ (2022) Subjective cognitive decline: Level of risk for future dementia and mild cognitive impairment, a meta-analysis of longitudinal studies. *Neuropsychol Rev* **32**, 703-735.
- [6] Wang X, Huang W, Su L, Xing Y, Jessen F, Sun Y, Shu N, Han Y (2020) Neuroimaging advances regarding subjective cognitive decline in preclinical Alzheimer's disease. *Mol Neurodegener* **15**, 55.
- [7] Amariglio RE, Mormino EC, Pietras AC, Marshall GA, Vannini P, Johnson KA, Sperling RA, Rentz DM (2015) Subjective cognitive concerns, amyloid- β , and neurodegeneration in clinically normal elderly. *Neurology* **85**, 56-62.
- [8] Perrotin A, La Joie R, de La Sayette V, Barré L, Mézenge F, Mutlu J, Guilleaume D, Egret S, Eustache F, Chételat G (2017) Subjective cognitive decline in cognitively normal elders from the community or from a memory clinic: Differential affective and imaging correlates. *Alzheimers Dement* **13**, 550-560.
- [9] Verberk IMW, Thijssen E, Koelewijn J, Mauroo K, Vanbrabant J, de Wilde A, Zwan MD, Verfaillie SCJ, Ossenkuppe R, Barkhof F, van Berckel BNM, Scheltens P, van der Flier WM, Stoops E, Vanderstichele HM, Teunissen CE (2020) Combination of plasma amyloid beta(1-42/1-40) and glial fibrillary acidic protein strongly associates with cerebral amyloid pathology. *Alzheimers Res Ther* **12**, 118.
- [10] Mielke MM, Hagen CE, Xu J, Chai X, Vemuri P, Lowe VJ, Airey DC, Knopman DS, Roberts RO, Machulda MM, Jack CR, Petersen RC, Dage JL (2018) Plasma phospho-tau181 increases with Alzheimer's disease clinical severity and is associated with tau- and amyloid-positron emission tomography. *Alzheimers Dement* **14**, 989-997.
- [11] Corriveau-Lecavalier N, Duchesne S, Gauthier S, Hudon C, Kergoat M-J, Mellah S, Belleville S (2020) A quadratic function of activation in individuals at risk of Alzheimer's disease. *Alzheimers Dement (Amst)* **12**, e12139.
- [12] Leal SL, Landau SM, Bell RK, Jagust WJ (2017) Hippocampal activation is associated with longitudinal amyloid accumulation and cognitive decline. *Elife* **6**, e22978.
- [13] Erk S, Spottke A, Meisen A, Wagner M, Walter H, Jessen F (2011) Evidence of neuronal compensation during episodic memory in subjective memory impairment. *Arch Gen Psychiatry* **68**, 845-852.
- [14] Rodda JE, Dannhauser TM, Cutinha DJ, Shergill SS, Walker Z (2009) Subjective cognitive impairment: Increased prefrontal cortex activation compared to controls during an encoding task. *Int J Geriatr Psychiatry* **24**, 865-874.
- [15] Pini L, Wennberg AM (2021) Structural imaging outcomes in subjective cognitive decline: Community vs. clinical-based samples. *Exp Gerontol* **145**, 111216.
- [16] Tromp D, Dufour A, Lithfous S, Pebayle T, Després O (2015) Episodic memory in normal aging and Alzheimer disease: Insights from imaging and behavioral studies. *Ageing Res Rev* **24**, Part B, 232-262.
- [17] Insel PS, Mormino EC, Aisen PS, Thompson WK, Donohue MC (2020) Neuroanatomical spread of amyloid β and tau in Alzheimer's disease: Implications for primary prevention. *Brain Commun* **2**, fcaa007.
- [18] Sperling RA, Dickerson BC, Pihlajamaki M, Vannini P, LaViolette PS, Vitolo OV, Hedden T, Becker JA, Rentz DM, Selkoe DJ, Johnson KA (2010) Functional alterations in memory networks in early Alzheimer's disease. *Neuro-molecular Med* **12**, 27-43.
- [19] Cabeza R (2002) *Hemispheric Asymmetry Reduction in Older Adults: The HAROLD Model*.
- [20] Wagner AD, Shannon BJ, Kahn I, Buckner RL (2005) Parietal lobe contributions to episodic memory retrieval. *Trends Cogn Sci* **9**, 445-453.
- [21] Sestieri C, Shulman GL, Corbetta M (2017) The contribution of the human posterior parietal cortex to episodic memory. *Nat Rev Neurosci* **18**, 183-192.
- [22] Edde M, Dilharreguy B, Theaud G, Chanraud S, Helmer C, Dartigues J-F, Amieva H, Allard M, Descoteaux M, Catheline G (2020) Age-related change in episodic memory: Role of functional and structural connectivity between the ventral posterior cingulate and the parietal cortex. *Brain Struct Funct* **225**, 2203-2218.
- [23] Mitolo M, Gardini S, Fasano F, Crisi G, Pelosi A, Pazzaglia F, Caffarra P (2013) Visuospatial memory and neuroimaging correlates in mild cognitive impairment. *J Alzheimers Dis* **35**, 75-90.
- [24] Fasano F, Mitolo M, Gardini S, Venneri A, Caffarra P, Pazzaglia F (2018) Combining structural magnetic resonance imaging and visuospatial tests to classify mild cognitive impairment. *Curr Alzheimer Res* **15**, 237-246.
- [25] Cona G, Scarpazza C (2019) Where is the "where" in the brain? A meta-analysis of neuroimaging studies on spatial cognition. *Hum Brain Mapp* **40**, 1867-1886.
- [26] Jacobs HIL, Boxtel MPJV, Heinecke A, Gronenschild EHB, Backes WH, Ramakers IHGB, Jolles J, Verhey FRJ (2012) Functional integration of parietal lobe activity in early Alzheimer disease. *Neurology* **78**, 352-360.
- [27] Gregory S, Long Jeffrey D, Tabrizi SJ, Rees G (2017) Measuring compensation in neurodegeneration using MRI. *Curr Opin Neurol* **30**, 380-387.
- [28] Brill E, Krebs C, Falkner M, Peter J, Henke K, Züst M, Minkova L, Brem A-K, Klöppel S (2022) Can a serious game-based cognitive training attenuate cognitive decline related to Alzheimer's disease? Protocol for a randomized controlled trial. *BMC Psychiatry* **22**, 552.

- [29] Molinuevo JL, Rabin LA, Amariglio R, Buckley R, Dubois B, Ellis KA, Ewers M, Hampel H, Klöppel S, Rami L, Reiberger B, Saykin AJ, Sikkes S, Smart CM, Snitz BE, Sperling R, Flier WM van der, Wagner M, Jessen F (2017) Implementation of subjective cognitive decline criteria in research studies. *Alzheimers Dement* **13**, 296-311.
- [30] Petersen RC (2016) Mild cognitive impairment. *Continuum (Minneapolis)* **22**, 404-418.
- [31] Carson N, Leach L, Murphy KJ (2018) A re-examination of Montreal Cognitive Assessment (MoCA) cutoff scores. *Int J Geriatr Psychiatry* **33**, 379-388.
- [32] Fazekas F, Barkhof F, Wahlund LO, Pantoni L, Erkinjuntti T, Scheltens P, Schmidt R (2002) CT and MRI rating of white matter lesions. *Cerebrovasc Dis* **13**, 31-36.
- [33] Nasreddine ZS, Phillips NA, Bédirian V, Charbonneau S, Whitehead V, Collin I, Cummings JL, Chertkow H (2005) The Montreal Cognitive Assessment, MoCA: A brief screening tool for mild cognitive impairment. *J Am Geriatr Soc* **53**, 695-699.
- [34] VLMT – Verbaler Lern- und Merkfähigkeitstest – Hogrefe Verlag, <https://www.testzentrale.ch/shop/verbaler-lern-und-merkfahigkeitstest.html>.
- [35] Rey A (1941) L'examen psychologique dans les cas d'encéphalopathie traumatique. (Les problems.). *Arch Psychol* **28**, 215-285.
- [36] Zelazo PD, Anderson JE, Richler J, Wallner-Allen K, Beaumont JL, Conway KP, Gershon R, Weintraub S (2014) NIH Toolbox Cognition Battery (CB): Validation of executive function measures in adults. *J Int Neuropsychol Soc* **20**, 620-629.
- [37] McKenna PAT, Warrington EK (1980) Testing for nominal dysphasia. *J Neurol Neurosurg Psychiatry* **43**, 781-788.
- [38] Guay F, Vallerand RJ, Blanchard C (2000) On the assessment of situational intrinsic and extrinsic motivation: The Situational Motivation Scale (SIMS). *Motiv Emot* **24**, 175-213.
- [39] Endicott J, Nee J, Harrison W, Blumenthal R (1993) Quality of Life Enjoyment and Satisfaction Questionnaire: A new measure. *Psychopharmacol Bull* **29**, 321-326.
- [40] Graf C (2008) The Lawton instrumental activities of daily living scale. *Am J Nurs* **108**, 52-62; quiz 62-63.
- [41] Oldfield RC (1971) The assessment and analysis of handedness: The Edinburgh inventory. *Neuropsychologia* **9**, 97-113.
- [42] Yesavage JA, Brink TL, Rose TL, Lum O, Huang V, Adey M, Leirer VO (1983) Development and validation of a geriatric depression screening scale: A preliminary report. *J Psychiatr Res* **17**, 37-49.
- [43] Baykara E, Kuhn C, Linz N, Tröger J, Karbach J (2022) Validation of a digital, tablet-based version of the Trail Making Test in the Delta platform. *Eur J Neurosci* **55**, 461-467.
- [44] Thijssen EH, Verberk IMW, Vanbrabant J, Koelewijn A, Heijst H, Scheltens P, van der Flier W, Vanderstichele H, Stoops E, Teunissen CE (2021) Highly specific and ultrasensitive plasma test detects Abeta(1-42) and Abeta(1-40) in Alzheimer's disease. *Sci Rep* **11**, 9736.
- [45] Klink K, Peter J, Wyss P, Klöppel S (2020) Transcranial electric current stimulation during associative memory encoding: Comparing tACS and tDCS effects in healthy aging. *Front Aging Neurosci* **12**, 66.
- [46] de Quervain DJ-F, Papassotiropoulos A (2006) Identification of a genetic cluster influencing memory performance and hippocampal activity in humans. *Proc Natl Acad Sci U S A* **103**, 4270-4274.
- [47] Wagner M, Wolf S, Reischies FM, Daerr M, Wolfsgruber S, Jessen F, Popp J, Maier W, Hull M, Frolich L, Hampel H, Pernecky R, Peters O, Jahn H, Luckhaus C, Gertz H-J, Schroder J, Pantel J, Lewczuk P, Kornhuber J, Wiltfang J (2012) Biomarker validation of a cued recall memory deficit in prodromal Alzheimer disease. *Neurology* **78**, 379-386.
- [48] Ebner NC, Riediger M, Lindenberger U (2010) FACES—A database of facial expressions in young, middle-aged, and older women and men: Development and validation. *Behav Res Methods* **42**, 351-362.
- [49] Seydell-Greenwald A, Ferrara K, Chambers CE, Newport EL, Landau B (2017) Bilateral parietal activations for complex visual-spatial functions: Evidence from a visual-spatial construction task. *Neuropsychologia* **106**, 194-206.
- [50] Suzuki A, Shinozaki J, Yazawa S, Ueki Y, Matsukawa N, Shimohama S, Nagamine T (2018) Establishing a new screening system for mild cognitive impairment and Alzheimer's disease with mental rotation tasks that evaluate visuospatial function. *J Alzheimers Dis* **61**, 1653-1665.
- [51] Penny WD, Friston KJ, Ashburner JT, Kiebel SJ, Nichols TE (2011) *Statistical parametric mapping: The analysis of functional brain images*, Elsevier.
- [52] RStudio Team (2015) *RStudio: integrated development for R*. RStudio Inc., Boston, MA.
- [53] Vannini P, O'Brien J, O'Keefe K, Pihlajamäki M, LaViolette P, Sperling RA (2011) What goes down must come up: Role of the posteromedial cortices in encoding and retrieval. *Cereb Cortex* **21**, 22-34.
- [54] Klöppel S, Vongerichten A, Eimeren T van, Frackowiak RSJ, Siebner HR (2007) Can left-handedness be switched? Insights from an early switch of handwriting. *J Neurosci* **27**, 7847-7853.
- [55] Klöppel S, Gregory S, Scheller E, Minkova L, Razi A, Durr A, Roos RAC, Leavitt BR, Papoutsis M, Landwehrmeyer GB, Reilmann R, Borowsky B, Johnson H, Mills JA, Owen G, Stout J, Scahill RI, Long JD, Rees G, Tabrizi SJ (2015) Compensation in preclinical Huntington's disease: Evidence from the Track-On HD Study. *EBioMedicine* **2**, 1420-1429.
- [56] Spets DS, Slotnick SD (2021) Are there sex differences in brain activity during long-term memory? A systematic review and fMRI activation likelihood estimation meta-analysis. *Cogn Neurosci* **12**, 163-173.
- [57] Verberk IMW, van Harten AC, Gouda M, Hussainali Z, van Engelen M-PE, Wilson D, Lemstra AW, Pijnenburg YAL, van der Flier WM, Teunissen CE (2022) Implementation of the Alzheimer's blood-based biomarker panel in clinical practice: The development of cutoffs for plasma Abeta1-42/1-40, P-tau181, GFAP and NfL across the clinical continuum. *Alzheimers Dement* **18**, e064069.
- [58] Bayoumy S, Verberk IMW, den Dulj B, Hussainali Z, Zwan M, van der Flier WM, Ashton NJ, Zetterberg H, Blennow K, Vanbrabant J, Stoops E, Vanmechelen E, Dage JL, Teunissen CE (2021) Clinical and analytical comparison of six Simoa assays for plasma P-tau isoforms P-tau181, P-tau217, and P-tau231. *Alzheimers Res Ther* **13**, 198.
- [59] Paulraj SR, Schendel K, Curran B, Dronkers NF, Baldo JV (2018) Role of the left hemisphere in visuospatial working memory. *J Neurolinguistics* **48**, 133-141.
- [60] Scheller E, Schumacher LV, Peter J, Lahr J, Wehrle J, Kaller CP, Gaser C, Klöppel S (2018) Brain aging and APOE epsilon4 interact to reveal potential neuronal compensation in healthy older adults. *Front Aging Neurosci* **10**, 74.
- [61] Corriveau-Lecavalier N, Rajah MN, Mellah S, Belleville S (2021) Latent patterns of task-related functional connectiv-

- 1329 ity in relation to regions of hyperactivation in individuals at
1330 risk of Alzheimer's disease. *Neuroimage Clin* **30**, 102643.
- 1331 [62] Li Y, Schindler SE, Bollinger JG, Ovod V, Mawuenyega
1332 KG, Weiner MW, Shaw LM, Masters CL, Fowler CJ, Tro-
1333 janowski JQ, Korecka M, Martins RN, Janelidze S, Hansson
1334 O, Bateman RJ (2022) Validation of plasma amyloid- β
1335 42/40 for detecting Alzheimer disease amyloid plaques.
1336 *Neurology* **98**, e688-e699.
- 1337 [63] Thijssen EH, La Joie R, Wolf A, Strom A, Wang P, Iaccarino
1338 L, Bourakova V, Cobigo Y, Heuer H, Spina S, VandeVrede
1339 L, Chai X, Proctor NK, Airey DC, Shcherbinin S, Duggan
1340 Evans C, Sims JR, Zetterberg H, Blennow K, Karydas AM,
1341 Teunissen CE, Kramer JH, Grinberg LT, Seeley WW, Rosen
1342 H, Boeve BF, Miller BL, Rabinovici GD, Dage JL, Rojas
1343 JC, Boxer AL (2020) Diagnostic value of plasma phospho-
1344 rylated tau181 in Alzheimer's disease and frontotemporal
1345 lobar degeneration. *Nat Med* **26**, 387-397.
- [64] Erhardt EB, Rachakonda S, Bedrick EJ, Allen EA, Adali 1346
 T, Calhoun VD (2011) Comparison of multi-subject ICA 1347
 methods for analysis of fMRI data. *Hum Brain Mapp* **32**, 1348
 2075-2095. 1349
- [65] Viviano RP, Damoiseaux JS (2020) Functional neuroimag- 1350
 ing in subjective cognitive decline: Current status and a 1351
 research path forward. *Alzheimers Res Ther* **12**, 23. 1352

Uncorrected Author Proof