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Abstract

Introduction: Certain personality traits are associated with higher risk of Alzheimer’s disease, similar to cognitive impairment. The identification of biological markers associated with personality in mild cognitive impairment could advance the early detection of Alzheimer’s disease.

Methods: We used hierarchical multivariate linear models to quantify the interaction between personality traits, state of cognitive impairment, and MRI biomarkers (gray matter brain volume, gray matter mean water diffusion) in the medial temporal lobe (MTL).

Results: Over and above a main effect of cognitive state, the multivariate linear model showed significant interaction between cognitive state and personality traits predicting MTL abnormality. The interaction effect was mainly driven by neuroticism and its facets (anxiety, depression, and stress) and was associated with right-left asymmetry and an anterior to posterior gradient in the MTL.

Discussion: Our results support the hypothesis that personality traits can alter the vulnerability and pathoplasticity of disease and therefore modulate related biomarker expression.

Keywords: Alzheimer’s disease; Brain; Personality; Neuroimaging; Psychiatry; Prognosis

1. Background

Translational research in Alzheimer’s disease (AD) has relied mostly on tests based on the assessment of cognitive state to detect individuals at risk, the individuals with mild cognitive impairment (MCI), and to identify the corresponding disease signatures such as medial temporal lobe (MTL) atrophy [1–3]. Looking beyond the unidimensional concept of MCI, current research aims to identify other important risk factors (genetic, personality traits) that would improve the prognostic accuracy of current tests and explain the high degree of individual variability in MTL atrophy that is not associated with cognitive decline.

In AD, personality changes, like cognitive decline, are also salient features of the disease [2–9]. In previous studies, interest in personality traits, particularly neuroticism (tendency to feel negative emotions such as stress, depression) and conscientiousness (tendency to be self-disciplined), and AD [10–13] was motivated by the fact that personality traits are stable into adulthood [14], have genetic-environmental underpinnings [15], brain anatomy correlates [16], and are also...
predictive of late-life developments such as cognitive dysfunction [7] or psychiatric symptoms [17]. However, the influence of personality traits on disease causation and their biological manifestations still remain unclear [6,8,18].

Our study aims to test whether morbid personality traits relate to individual differences within the MTL independent of cognitive state. To further improve the discrimination between the two groups in term of brain anatomical changes, we aim to identify the personality profile that minimizes the variance within each group (MCI and non-cognitively impaired [NCI]) and maximizes the difference between them. We predict that neuroticism and its underlying facets, anxiety, depression, and stress will have the greatest exacerbating effects on disease stages. We also expect that identified personality profile will correlate with known functional organization within the MTL.

To quantify MTL atrophy, we used structural magnetic resonance imaging and derived measures of gray matter volume (GMV) and gray matter mean diffusivity (GMMD). GMMD is considered a more sensitive marker than GMV to detect MTL abnormalities in MCI [19]. We used a multivariate strategy [20] to provide a comprehensive test of the association between personality traits, cognitive state, and brain anatomy. The method (Fig. 1) is data driven, unbiased, takes into account the multidimensional and hierarchical nature of the five-factor model of personality, and uses anatomical constraints to decompose the different sources of variability.

2. Methods

2.1. Participants

The study included older adults selected from a longitudinal cohort recruited in the psychogeriatric and geriatric memory clinics of the Lausanne University hospital. The local ethics committee gave permission for the research protocol, and all participants gave written informed consent before taking part in the study. All participants completed comprehensive clinical, psychiatric, and cognitive assessments at the time of MRI scanning. Participants with psychiatric or neurological central nervous system disorders (stroke, tumor), dementia, and alcohol or drug abuse were excluded. The 97 participants included in the study were divided into two groups, MCI and NCI, according to the conventional Winblad criteria [2] where MCI is defined as abnormal but does not fulfill the diagnostic criteria for dementia. A total of 29 participants were MCI (8 males, aged 68 ± 8 years, Mini–Mental State Examination [21] [MMSE]: 27.7 ± 1/range [25–29], Clinical Dementia Rating [22] [CDR] = 0.5, with MCI amnestic 23, nonamnestic 6) and 68 were NCI (18 males, aged 66 ± 6 years, MMSE: 29.1 ± 1/range [26–30], CDR = 0).

2.2. Personality and neuropsychological/psychiatric assessments

To obtain reliable measures of current personality profiles, we asked relatives of participants to complete the

Fig. 1. (A) NEO Personality inventory (NEO-Pi-R) is hierarchical construct composed of 5 domains and 6 facets for each domain. (B) Search volume of interest with the hippocampus in yellow and parahippocampal cortex in red. (C) Multivariate Linear Model (MLM) identified the personality profile and the brain distributed pattern that best explain the covariance between personality scores and anatomical measures.
240-item NEO Personality Inventory Revised (NEO-PI-R) [23]. This questionnaire, rated on a five-point agreement scale, is based on the five-factor model of personality derived from statistical factor analysis of various personality lexical inventories [23]. It is hierarchically divided into five broad domains (Fig. 1A): neuroticism (a tendency to feel negative affects and to be susceptible to psychological distress), extraversion (a tendency to be sociable and lively), openness (a tendency to be open to new experiences), agreeableness (a tendency to be cooperative, altruistic, and trusting), and conscientiousness (a tendency to be careful, dutiful, and responsible). Each domain contains six facets (Fig. 1A). The facets of the neuroticism domain are anxiety, anger, hostility, depression, self-consciousness, impulsiveness, and vulnerability to stress. The NEO-PI-R has a high test-retest reliability in the elderly [23] and high inter-rater reliability in patients with AD [24].

Internal reliability of NEO-PI-R scores was estimated with Cronbach’s alpha. In our sample, the values ranged from 0.63 to 0.68 for the NEO-PI-R domains and from 0.79 to 0.87 for the facets of neuroticism (a nominal value of 0.7 denotes internal consistency [25]). Other rating scales and tests used included the Hospital Anxiety and Depression Scale (HADS-A [anxiety] and HADS-D [depression], respectively) [26] and the RL-RI-48 memory item task [27].

2.3. Neuroimaging data

Data was acquired using whole-brain MRI T1-weighted (T1w) structural images (structural magnetic resonance imaging protocol: 1-mm isotropic resolution with a matrix of 256×256 voxels, repetition time 2.3 seconds, echo time 2.91 seconds) and diffusion-weighted MRI (1.8 mm³ resolution, with a matrix of 128×128 voxels, 30 directions, high b-value of 1000 seconds/mm²) on a 3T MRI scanner (Siemens Trio).

We applied a standard data preprocessing pipeline using a statistical parametric mapping package [28,29] (SPM8-Matlab toolbox, www.fil.ion.ucl.ac.uk/spm) to the T1w images with a bias field correction and unified segmentation into white and gray matter tissue classes. In addition, we applied a standard preprocessing pipeline using FreeSurfer software [30] (FSL; http://www.fmrib.ox.ac.uk/fsl/) with correction for eddy currents and head movement distortion before extracting mean diffusion images. Mean diffusion and T1w images were spatially realigned and normalized to Montreal Neurological Institute space with the DARTEL procedure contained in the Voxel-Based Quantification toolbox [31]. The final outputs were restricted to the gray matter segment to obtain voxel-wise estimations of GMV and GMMMD. Finally, we smoothed the images with an isotropic gaussian kernel of 8 mm full width at half maximum. Anatomical labeling was based on the Automated Anatomical Labeling atlas [32].

2.4. Statistical analyses

2.4.1. Anatomical differences explained by cognitive state stratification

We first conducted a univariate regression analysis to test for differences in the brain measures (GMV and GMMMD) between MCI and NCI groups. The model included the cognitive state stratification factor, age, and total intracranial volume as confounding variables.

2.4.2. Anatomical differences explained by personality domains and cognitive state

Secondly, we used a multivariate model [20] to address the question whether, beyond cognitive factors, there are specific personality traits that could explain anatomical MTL differences between MCI and NCI. In the literature, multivariate factorial analysis has often been used in studies of personality to extract significant factorial structures [23,33,34]. We used a variant of this method, the multivariate linear method (MLM) that is similar to standard multivariate factorial analysis but additionally integrates anatomical information with the cognitive variables and any confounds. The MLM procedure is based on singular value decomposition, which summarizes covariance between personality scores (Fig. 1A) and anatomical data (Fig. 1B). The output of the MLM (Fig. 1C) consists of pairs of spatially distributed brain patterns associated with a set of linear combinations of personality traits that are maximally correlated with them. The significance of the personality profiles is assessed by a multivariate F-test (based on partial averages of the eigenvalues) that defines the spaces of interest for the five personality domains beyond those of the cognitive and other confounding factors. Post hoc univariate analyses were then performed with identified profiles to determine their mapping at the voxel level.

2.4.3. Anatomical differences explained by personality facets and cognitive state

We performed the MLM analysis at the facet level within the whole search volume of interest.

3. Results

3.1. Demographic, personality traits, and neuropsychological/psychiatric results

We found (see details in Table 1) no statistical differences between MCI and NCI groups for all demographic variables (age, gender). As expected, the MMSE was significantly different between the two groups, and the memory scores measured by the RL-RI-48 memory item task were significantly lower in the MCI group. There were no statistical differences in HADS-D for depressive symptoms and HADS-A for anxiety symptoms scores. Neuroticism scores were also significantly higher in MCI patients (Table 1).
Demographic variables and neuropsychological scores

| Demographic variables | Non-cognitively impaired (mean ± SD) | MCI (mean ± SD) | T- or χ²-statistic (d.o.f: degree) | P value |
|-----------------------|-------------------------------------|---------------|---------------------------------|--------|
| n                     | 68                                  | 29            |                                 |        |
| CDR                   | 0                                   | 0.5           |                                 |        |
| MMSE                  | 29.1 ± 1                            | 27.7 ± 1      | 5.4 (95)                        | 0      |
| Age                   | 66 ± 6                              | 68 ± 8        | -1.6 (95)                       | .1     |
| Gender (Female/Male)  | 0.7                                 | 0.7           | 0.01 (95)                       | .9     |
| Education level       | 2                                   | 2             | 0.4 (2)                         | .8     |

Personality: domain scores (NEO PI-R)

- Neuroticism: 77.6 ± 23
- Extraversion: 105.7 ± 18
- Openness: 109.9 ± 19
- Agreeableness: 135.2 ± 17
- Conscientiousness: 134.5 ± 19

Other neuropsychological scores

- Cued recall (RL-RI-48): 29.2 ± 4
- Depression score (HADS-D): 2.1 ± 2
- Anxiety score (HADS-A): 4.7 ± 3

Abbreviations: CDR, Clinical Dementia Rating; HADS, Hospital Anxiety and Depression Scale; MCI, mild cognitive impairment; MMSE, Mini-Mental State Examination; NEO PI-R, NEO Personality Inventory Revised.

3.2. Anatomical differences related to cognitive state stratification

Using a whole-brain familywise error correction, we found no significant differences in GMV. However, with the same stringent level of correction for multiple comparisons, GMMD was significantly different between the two groups in several regions. In the MTL, local maxima for these differences (Table 2) were located in both parahippocampal and hippocampal subregions (cornu ammonis, dentate gyrus, and subiculum, according to the probabilistic cytoarchitectonic map [35]). At the whole-brain level, GMMD differences were significant in the left middle temporal cortex \((Z = 5.06, \text{xyz} = [-57.13.5.3])\), right superior temporal cortex \((Z = 4.74, \text{xyz} = [54.0.3])\), and left insula \((Z = 4.84, \text{xyz} = [-41.12.14])\). The right temporal pole \((Z = 4.83, \text{xyz} = [-41.3.9])\), and left lingual cortex \((Z = 4.65, \text{xyz} = [-12.36.0])\), and right postcentral gyrus \((Z = 4.79, \text{xyz} = [53.24.56])\). Inclusion of education level and gender did not add contributive information to the brain measures.

3.3. Anatomical differences related to personality domains and cognitive state

The multivariate analysis of GMV showed that there was a significant contribution of personality domains to alterations in brain structure. The first component identified related to personality traits (Fig. 2A) was significant \((F = 3.77, P < e-5)\) and explained 54.39% of the between-group covariance in the search volume of interest (Fig. 2B). Neuroticism and agreeableness were identified as the main domains contributing to this brain component (Fig. 2A). No other regions showed significant differences between the two groups.

Post hoc univariate regression analyses of GMV were performed with the first component of the MLM analysis as predictor with domain level. This revealed significant structural differences located in both parahippocampal cortices (in the entorhinal cortex and subiculum) (Table 2).

The MLM analysis of GMMD also showed a significant contribution of personality traits (Fig. 2C) to the first component \((F = 5.32, P < .001)\), which explained 69.24% of the between-group covariance in the search volume of interest (Fig. 2D). The neuroticism and agreeableness domains had more weight than the three others (Fig. 2C), and a distributed spatial pattern of brain differences was revealed in the right hippocampal and parahippocampal cortices (Fig. 2D).

3.4. Anatomical differences related to personality facets of neuroticism and cognitive state

The MLM analysis of GMV showed contributions from the neuroticism facets profile (Fig. 2E) in the MTL region, mainly in the right hemisphere (Fig. 2F). The first
component was significant ($F = 8.84, P = 0$) and explained 72.71% of the covariance (Fig. 2E and F).

The MLM analysis of GMMD again revealed a significant contribution of the personality facets profile (Fig. 2G), with the first component significant ($F = 3.52, P < .0005$) and explaining 46.72% of the variance in the MTL, mainly in the anterior part (Fig. 2H).

4. Discussion

Our multifactorial and multivariate analysis decomposes the complex relationship between three risk markers of AD, namely (1) anatomical atrophy, (2) cognitive decline, and (3) personality traits, which together reveal clinical and topographical signatures in MCI that has direct implications for refining current models of AD.

Our results are important because, with a few exceptions, there is a paucity of data-linking personality to neurobiological mechanisms of disease. A few neuropathologic studies [8,11,13,36] of confirmed AD cases have provided evidence for a role of higher levels of neuroticism and depression in relation to disease symptoms (i.e., a more rapid cognitive decline in old age and higher risk of AD [11]) but provide ambiguous evidence for a direct link with lesions observed at autopsy (no link or linked with a higher level of neurofibrillary tangles and neuritic plaques) [8,11,36]. Neuroimaging studies of personality have been mainly conducted in healthy adults and have found significant associations between neuroticism trait and structural differences in frontal and temporal regions [16]. A recent study showed that in MCI individuals, the severity of white matter lesions in the MTL, but not the atrophy, was associated with higher neuroticism and lower conscientiousness [18]. Here, we investigate the multivariate relationship between personality and the MTL and provide a link to the vast majority of AD neuroimaging studies that report consistent effects of stress and depressive symptoms or cognitive and AD states on the hippocampus [37,38].
We identified a spatial pattern of anatomical alterations in MCI individuals compared with NCI that extends beyond the temporal cortex to the left insula, the left lingual cortex, and the right postcentral gyrus for GMMD. We also observed that diffusion-based measures are more sensitive than volumetric ones to detect brain abnormality in MCI, which is in line with recent findings [19]. GMMD differences may be caused by modifications in the intra/extracellular space due to pre-atrophic changes.

We also identified specific anatomical patterns associated with personality traits. The MLM analysis of GMMD with personality domains revealed an asymmetry between the right and left MTL. The anatomical changes associated with the facet level of neuroticism showed a spatial gradient from the anterior to posterior parts of the MTL. For GMV this asymmetry was also observed at the facet level of neuroticism. Other studies on the effect of stress and depression on healthy and depressive individuals have also reported differences between the left and right hippocampal areas that could be explained by neurochemical and brain tissue property differences [39,40]. The anteroposterior gradient has been related to the specific role of the anterior hippocampus in stress and emotion-related behavior [41]. The link between depression and AD has also been clinically reported, with chronic distress and depression being risk factors for cognitive decline in AD [7,42]. Our results, in light of these studies, converge to suggest that depression and AD share biological substrates in the hippocampus that are stress related [43,44].

Our findings highlight neuroticism, agreeableness and facets of anxiety, stress, hostility, and depression as key the explanatory variables of anatomical changes in the...
MTL. High scores for neuroticism are often reported to be predictive of cognitive impairment in AD [6,7,11,12]. This is related to the occurrence of neuropsychiatric problems (e.g., depression and anxiety symptoms), higher comorbidity of mental disorders, lower quality of life, and shorter life expectancy [17,45]. A premorbid agreeableness is linked to agitation and irritability in AD [4,46]. The low level of conscientiousness also predicts conversion of MCI to AD [13]. Although personality traits such as openness and extraversion [4,5] have been associated with early AD and MCI, we found less effect of these domains in our study. This can be due to fact that in our study, we identified the personality traits that correlate with brain changes and not just cognition stratification.

Our data suggest that personality is an important parameter that needs to be included in the modeling of the pathophysiological processes leading to AD [7,47]. Recently, a new model has been proposed by Jack et al. (2013) [47] in which different states of the AD biomarkers (e.g., brain atrophy, tau, amyloid β, memory, and clinical function) each follow a sigmoid-shaped curve. The authors argue that for AD model, the most informative parameters are the onsets of curves on the horizontal time axis, their slopes, and their temporal ordering. We suggest adding another level to this model based on the psychopathology literature, whereby a specific personality trait, such as proneness to stress or depression, can affect the shape and the temporal ordering of biomarker state curves by two main mechanisms (Fig. 3). The first mechanism is predisposition/vulnerability, where a certain personality trait profile modulates the risk of disease and changes the onset of biomarker curves. The second exacerbating mechanism is pathoplasticity, where a personality trait has an additive (or diminishing) effect on the course of disease and hence impacts the slopes of the temporal curves.

Therefore, other features, such as behavioral and psychological symptoms [45], pathophysiological markers of AD, genetic susceptibility factors such as serotonin [48,49], and personality, as demonstrated here and elsewhere, may help to clarify mechanisms to explain the associations between cognitive decline and MTL atrophy in aging.

Acknowledgments

A.v.G. is supported by a Swiss National Science Foundation FNS 3200BO-122263. B.D. is supported by a Swiss National Science Foundation (NCCR Synapsy, project grant Nr 320030_135679 and SPUM 33CM30_140332/1), Foundation Parkinson Switzerland, Foundation Synapsis, the Novartis Foundation for medical biological research, and the German Research Council (DFG, KFO 247). Laboratoire de Recherche en Neuroimagerie is supported by generous funding from the Roger de Spoelbergh and Partridge foundations. F.K. received funding from the European Union Seventh Framework Programme (FP7/2007-2013) under grant agreement number 604102 (HBP Ramp-Up Phase) and grant agreement number 720270 (HBP SGA1), the VELUX STIFTUNG and Pharnext, Paris. The authors have declared that no conflict of interest exists.

RESEARCH IN CONTEXT

1. Systematic review: Most prior studies have focused on personality changes after Alzheimer’s disease onset. Fewer studies reported associations between postmortem pathological markers and personality traits. A majority of studies examined the effect of one or two traits separately to dementia incidence and also reported significant contribution of neuroticism.

2. Interpretation: We found a specific association between cognitive decline, personality traits, and biological changes in the medial temporal lobe. Interestingly, from disease modeling point of view, the association was explained by a low-dimensional factor, which corresponded to a personality profile dominated by neuroticism trait associated with a spatial gradient along the medial temporal lobe.

3. Future directions: These findings add to the body of evidence that personality traits are stable features, which in the context of cognitive decline, lead to brain vulnerability. Multifactorial models, as proposed here, are needed to combine the different states and trait markers for accurate Alzheimer’s disease risk prediction.

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