

Genetic evidence for protective effects of angiotensin converting enzyme against Alzheimer's disease but not other neurodegenerative diseases in European populations

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Abstract

Background and Objectives:

Angiotensin converting enzyme (ACE) inhibitors are a commonly prescribed class of medication used to treat heart failure, hypertension and chronic kidney disease. However, previous observational studies have shown conflicting directions of associations between ACE inhibitors and risk of Alzheimer's disease. Genetic evidence has supported a protective effect of cerebral ACE against Alzheimer's disease (AD). However, it is unclear whether this effect is mediated through blood pressure and extends to other neurodegenerative diseases.

Methods:

We performed genetic colocalization investigating an effect of cortical *ACE* expression on AD risk in people of European ancestry. We further investigated whether any effect of *ACE* expression on AD risk is mediated through changes in blood pressure, and whether effects extend to Parkinson's disease, small vessel disease or cognitive function in a Mendelian randomization paradigm.

Results:

There was genetic evidence supporting a protective effect of cortical *ACE* expression on AD risk in people of European ancestry. Although higher cortical *ACE* expression was associated with higher blood pressure, there was no strong evidence to support that its association with AD was mediated through blood pressure, nor that *ACE* expression affected risk of other neurodegenerative traits.

Discussion:

Genetic evidence supports protective effects of cerebral *ACE* expression on AD, but not other neurodegenerative outcomes in people of European ancestry. Further work is required to investigate whether therapeutic inhibition of *ACE* increases risk of Alzheimer's disease.

Introduction

Alzheimer's disease (AD) is a leading cause of morbidity worldwide and its prevalence is projected to increase in line with the aging global population¹. Several pre-clinical and observational studies have implicated the role of central nervous system angiotensin converting enzyme (ACE) levels in the pathogenesis of AD. Cerebral ACE, and downstream product angiotensin II, are increased in patients with AD and promote neuroinflammatory cytokines, reduce acetylcholine release and attenuate cerebral blood flow – all factors implicated in the development of AD². Animal models have shown that hypertensive rats treated with centrally acting ACE inhibitors (e.g. captopril, perindopril), but not hydralazine, have significantly lower age-related impairment in learning and memory, regardless of changes in blood pressure³. Observational data also supports the neuroprotective role of central-acting ACE-inhibitors compared to predominantly peripherally acting ACE-inhibitors^{2,4}.

On the contrary, there is also some evidence that ACE may serve in preventing AD. For example, *in vitro* studies have supported that ACE degrades amyloid- β plaques, a pathological hallmark of AD⁵. Animal AD models with heterozygous deletion of the *ACE* gene demonstrated that a decrease in ACE levels promoted amyloid- β deposition and increased the number of apoptotic neurons⁴. At present, there is therefore uncertainty surrounding the role of ACE-inhibitors in the pathogenesis of AD.

Genetic data, in the form of genome-wide association studies (GWAS) can be leveraged in a paradigm known as Mendelian randomization (MR) to study the causal effect of an exposure on an outcome. Mendelian randomization (MR) utilises genetic polymorphisms to proxy the exposure in question and has notable benefits including the reduction in confounding and elimination of reverse causality. This occurs due to the random allocation of genetic variants and balancing of confounding factors at conception. It also enables us to study the effects of chronic changes in exposure on life-time risk of a disease, which is advantageous in the context of neurodegenerative traits, which oftentimes develop over many years. This form of study design makes several assumptions: the genetic proxy must be associated with the exposure; the genetic variant only affects the outcome through the exposure of interest with no horizontal pleiotropic effect and the genetic variant is not associated with any known confounder affecting the exposure and the outcome.

Recent GWAS have identified the *ACE* gene as a locus of interest in the development of AD⁶. Bivariate GWAS and colocalisation studies suggest that the *ACE* gene may mediate an association between blood pressure traits and AD risk, with the allele associated with lower SBP also associated with higher AD risk. Tissue-specific expression has demonstrated that higher cerebellar *ACE* expression has a positive association with AD risk⁷. By extension, this implicates a possible detrimental effect of centrally acting pharmacological ACE inhibition on AD risk. This is of direct clinical relevance as ACE inhibitors are one of the most commonly prescribed anti-hypertensive agents and are oftentimes commenced as a first-line medication in younger patients. Therefore, it is imperative to understand any potential long-term effect of ACE modulation on risk of later life neurodegenerative diseases.

To further explore the relationship between ACE and neurodegenerative diseases, we advance previous work by performing three-way colocalization analyses for *ACE* gene expression in the cortex, systolic blood pressure (SBP) and AD risk. This approach enables us to identify a genetic proxy for the effect of ACE inhibitors that cross the blood-brain barrier, and explore its association with risk of AD and other neurodegenerative diseases in a MR paradigm. Finally, we assess whether any association could be mediated by effects of SBP on AD risk. In this way, we elucidate the complex interplay between ACE and AD, and assess the potential effect of ACE-inhibition on neurodegenerative disease risk more widely.

Methods

Data sources

For both colocalization and Mendelian randomization (MR) analyses, data were obtained from publicly available summary statistics of genome-wide association studies (GWAS) outlined in Table 1. We selected Parkinson's disease, cognitive performance, lacunar stroke and MRI quantified white matter hyperintensity as outcomes to represent common neurodegenerative conditions. Lacunar stroke and MRI-quantified white matter hyperintensity, both which are well-recognised features of vascular dementia, were used as surrogate outcomes in the absence of available GWAS for vascular dementia. Finally, cognitive function was selected to proxy the effects of cortical ACE expression on disease-agnostic cognitive function. Cognition outcome data was obtained from a meta-analysis of the COGENT consortium GWAS and UK Biobank. This study defined cognition using a general cognitive function 'g', calculated based on the first unrotated component extracted

from a principal component analysis of individual test scores across a range of neuropsychological tests⁸. The largest publicly available GWAS for each trait at the time of the study was selected for inclusion.

Statistical analysis

We conducted colocalization analysis of genetic associations for *ACE* gene expression in brain cortex tissue within +/-20kb of the *ACE* gene (hg19 co-ordinates: chr17:61,554,422-61,599,205) and AD liability. In colocalization studies, we assess for the probability of a common shared variant between traits, which implies a likely shared causal mechanism. For pairwise colocalization between traits, we employed coloc,⁹ a Bayesian method to calculate posterior probabilities (PP) for the competing models:

1. Model 0: The genomic region does not contain a variant influencing the exposure or outcome.
2. Model 1: The genomic region contains a variant influencing only the exposure.
3. Model 2: The genomic region contains a variant influencing only the outcome.
4. Model 3: The genomic region contains two separate variants, one influencing the exposure and the other influencing the outcome.
5. Model 4: The genomic region contains a variant influencing both exposure and outcome.

The prior probability of any random variant being associated with both traits was set at 1×10^{-5} . We defined colocalization between the exposure and outcome if the PP is greater than 0.8 for model 4.

We also assessed for three-way colocalization with SBP using the HyPrColoc method¹⁰ (Hypothesis Prioritisation for multi-trait Colocalization), to investigate whereby AD risk, cortical *ACE* expression and blood pressure share a common causal variant. HyPrColoc estimates the posterior probability of full colocalization, that is, the probability of all traits sharing the same causal variant.¹⁰ The prior probability parameters in HyPrColoc were also set at their default values, i.e. the prior probability for a variant influencing one trait = 1×10^{-4} , and the conditional probability of a variant influencing another trait = 0.02.¹⁰

The single-nucleotide polymorphism (SNP) with the greatest PP from colocalization analysis of cortical *ACE* expression and AD risk represents the genetic proxy most likely to simulate the effect of cerebral cortex ACE modulation on AD risk. We then employ a MR paradigm to explore whether this variant is associated with other neurodegenerative traits.

MR employs genetic polymorphisms as instrumental variables to investigate the effects of an exposure on an outcome.¹¹ For consistent causal estimates, the genetic proxy must (i) be associated with the exposure, (ii) only affect the outcome through the exposure of interest and (iii) not associate with any confounder of the exposure and outcome.¹¹

Finally, two-sample MR was performed to investigate whether changes in blood pressure are responsible for mediating risk of AD and other neurodegenerative traits. Instruments to proxy systolic blood pressure (SBP) were selected as uncorrelated ($r^2 < 0.0001$) variants that significantly associated with SBP ($p < 5 \times 10^{-8}$). Variants within the *ACE* gene were not removed for this as the aim of the analysis was to look at generic reduction in SBP, regardless of the gene of interest. Odds ratios were derived using inverse-variance weighted pooling of individual SNP Wald ratios, which corresponds to a weighted regression with the precisions of the variant-outcome associations acting as weights, and the intercept fixed to zero. Sensitivity analysis was conducted to assess for potential pleiotropic effects using alternative pooling techniques (simple median, weighted median and MR-Egger). Median estimates are robust even in situations where up to 50% of the weights contributing to the analysis are from invalid instruments¹². MR-Egger is a pooling technique where the intercept in the weighted regression is not fixed to zero. The intercept term is used to indicate the average pleiotropic effects of the variants used¹². A non-significant intercept term suggests no evidence for unbalanced pleiotropic effects. This method is robust even when all instrumental variables are invalid, as long as the INstrument Strength Independent of Direct Effect (INSIDE) assumption holds: that any pleiotropic effect of the variants on the outcome are independent of the strength of their association with the exposure¹². All data analyses were performed using R statistical software, with “coloc” package version 4.1.0, “hyprcoloc” package version 1.0 and “TwoSampleMR” package version 4.26.

Standard protocol approvals, registrations, and patient consents

All data used for this study are publicly available and their original studies are cited. All these studies obtained relevant participant consent and ethical approval. No formal ethical approval was required for use of publicly available genome-wide association data.

Data availability statement

All data used in this study are publicly available. The statistical code used in this work is available from the corresponding author upon reasonable request.

Results

Genetic colocalization

Colocalization analysis provided evidence for a shared causal variant for each pairwise combination (PP for colocalization of cortical *ACE* gene expression and AD liability = 0.98; PP for colocalization of cortical *ACE* expression and SBP = 0.97; PP for SBP and AD liability = 0.98). In three-way colocalization by HyPrColoc, the estimated PP of full colocalization = 0.83. The variant rs4291 was the most likely shared causal variant for all traits. Similar results were obtained when only considering AD cases and not also AD-by-proxy (Supplementary eResults).

Mendelian randomization

The A allele of rs4291 that associated with lower cortical *ACE* expression was negatively associated with SBP (effect estimate per increase -0.28mmHg, 95% confidence interval -0.35 to -0.22). This same variant was positively associated with AD risk (odds ratio [OR] 1.06, 95% confidence interval [CI] 1.04 to 1.08). However, strong associations were not identified for the other neurodegenerative traits (OR for lacunar stroke 0.97 (95% CI 0.93 to 1.01); OR for Parkinson's disease 0.99 (95% CI 0.96 to 1.03); beta estimate for cognitive function 0.01 (95% CI -0.001 to 0.01); beta estimate for white matter hyperintensity -0.02 (95% CI -0.11 to 0.07; Supplementary eTable 2). Genetically predicted SBP was not associated with AD risk (OR 1.01, 95% confidence interval 1.00 to 1.01 per mmHg increase in SBP) using a genome-wide instrument, suggesting that while SBP colocalizes with cortical *ACE* expression at the *ACE* gene, reduction in SBP more generally is not directly associated with AD risk. These findings was consistent in sensitivity analysis (Supplementary eTable 3).

Discussion

This study leveraged genetic data to identify support for ACE in prevention of AD, with no strong evidence identified supporting effects of ACE on other neurodegenerative traits. While increased cortical *ACE* expression associated with lower AD risk, there was no MR evidence supporting that genetically predicted SBP affects risk of AD.

From a mechanistic perspective, ACE has been shown to breakdown neurotoxic amyloid-beta isoform (A β 42) to a less toxic form (A β 40). Administration of a clinical dose of ACE inhibitor to human amyloid precursor protein transgenic mice was associated with increased brain amyloid deposition⁴. In humans, patients with AD have lower A β 42-to-A β 40-converting activity compared with sera from age-matched healthy individuals⁴. Our current findings support that ACE protects against AD, although further work is required to investigate whether this is attributable to reduced amyloid aggregation or other unrelated mechanisms.

An observational study among 406 participants with mild-to-moderate AD demonstrated a reduction in cognitive decline for people receiving a centrally-acting ACE inhibitor (perindopril) compared to peripherally-acting ACE inhibitor¹⁹. Other studies have shown increased risk of incident dementia and disability associated with peripherally-acting ACE inhibitors compared to other anti-hypertensive medication²⁰. Conflicting findings between genetic and observational studies could be explained by MR being less liable to environmental confounding and reverse causality¹¹, due to the random allocation of genetic variants at conception.

Our current work is consistent with other genetic studies supporting a role of ACE in preventing AD and has several additional strengths. Firstly, obtaining association estimates from the MetaBrain consortium (n = 6,601 participants)¹³, we investigate cortical *ACE* expression and AD risk. This dataset is significantly larger than the GTEx resource (n = 205) that has been utilised in previous work⁷. Secondly, we investigate whether SBP mediates the relationship between ACE and AD risk, and do not find evidence that supports this. Finally, we explore the associations of genetically proxied cortical *ACE* expression with other traits and do not find evidence to support that this association applies across other neurodegenerative traits.

This work also has several limitations. Clinical diagnosis of AD is challenging as there is significant overlap in symptoms with other forms of dementia, limiting the specificity of case definitions in GWAS. To explore for this, we also assessed several other neurocognitive traits. Given the absence of strong evidence of ACE effects for these outcomes, it seems likely that our findings are specific for AD risk, rather than a generic effect on dementia or cognition. As with all studies leveraging genetic data, there remains the possibility of biological pleiotropy introducing confounding. It is also not possible to extrapolate the magnitude of clinical effect or required drug exposure for ACE inhibitors to represent a real-world risk for AD. Factors such as the ability of an ACE inhibitor to cross the blood-brain-barrier may also shape AD risk and should be further studied. Furthermore, our work was based on data obtained from individuals of European genetic ancestry, and it is unclear whether these findings extend to other ethnic groups. A strength of the current study is that we assess for association between cortical ACE expression and various neurodegenerative traits. The outcome GWAS data are obtained from relatively homogenous populations, with most studies utilising UK Biobank data. The consideration of participants of European genetic ancestry avoids bias related to combining results from different ancestry groups.

In summary, while ACE inhibitors have numerous indications and are the cornerstone of hypertension, chronic kidney disease and heart failure management, this study finds evidence for a beneficial effect of cerebral cortex ACE in preventing AD. It would be premature to alter current clinical practice based on this evidence, and rather these findings should encourage further research into the effect of ACE inhibitors on AD risk.

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Table 1: Sources for genetic data

Data	Data source	Population	Study overview	Adjustments
Cortex <i>ACE</i> gene expression	MetaBrain cortical eQTL GWAS 2021 ¹⁰	6,518 individuals of European ancestry	Human brain cortex RNA-seq datasets were obtained from 13 different studies.	Technical covariates.
Systolic blood pressure	Evangelou et al 2018 ¹¹	757,601 individuals of European ancestry	Systolic blood pressure defined as the mean of two automated values. Correction was made for anti-hypertensive medication use by adding 15 mmHg to the systolic blood pressure of participants receiving medication	Body mass index, sex, age and age ² ; study-specific covariates to adjust for technical variation and population stratification.
Alzheimer's disease	De Rojas et al 2020 ¹³	GWAS meta-analysis of up to (i) 36,675 cases and 58,482 controls, and Alzheimer's disease-by-proxy analysis of (ii) 27,696 cases of maternal Alzheimer's disease with 260,980 controls, and (iii) 14,338 cases of paternal Alzheimer's disease with 245,941 controls, all of European ancestry.	Case defined using clinical criteria (Diagnostic and Statistical Manual of Mental Disorders IV and the National Institute on Aging and Alzheimer's Association). Proxy cases defined on the presence of parental diagnosis of dementia.	Studies in part (i) were adjusted for first four genetic principal components (PCs). Studies in part (ii) and (iii) were adjusted for genetic PCs, age and sex.
Parkinson's disease	Nalls et al 2019 ¹⁴	GWAS meta-analysis of 37,688 cases and 18,618 UK Biobank proxy-cases (first-degree relative with Parkinson's disease) and 1.4 million controls.	Cases defined based on clinic visit and UK Brain Bank criteria for Parkinson's disease. Proxy cases within UK Biobank defined by self-reported family history of Parkinson's disease	Age at onset in cases (age at most recent examination for controls), biological sex and up to the first five genetic PCs
Lacunar stroke	Traylor et al 2021 ¹⁵	GWAS of 6,030 Cases, 219,389 controls in individuals of European ancestry	Cases defined using Trial of Org 10172 in Acute Stroke Treatment (TOAST) criteria for small vessel disease and/or MRI evidence of lacunar stroke	Genetic PCs.
White matter hyperintensity	Persyn et al 2020 ¹⁶	GWAS of 17,663 individuals of European ancestry in the UK Biobank	White matter hyperintensities quantified using fractional anisotropy on MRI brain.	Age, sex, assessment centre, MRI head motion indicators and genetic PCs.

Cognitive function	Lee et al 2018 ¹⁷	Meta-analysis of UK Biobank (n = 222,543) and the COGENT consortium (n = 35,298). All participants of European ancestry.	Cognition was defined in the UK Biobank as mean standardised score on a test of verbal-numerical reasoning. For the COGENT consortium, phenotype defined as the first principal component derived from three or more neuropsychological tests.	Age, sex, interaction for sex and age, first ten genetic PCs.
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Table 1: Details genome-wide association studies used in the present study. COGENT consortium: Cognitive Genomic Consortium, eQTL: expression quantitative trait loci, GWAS: genome-wide association study, MRI: magnetic resonance imaging, PC: principal component, RNA-seq: RNA sequencing

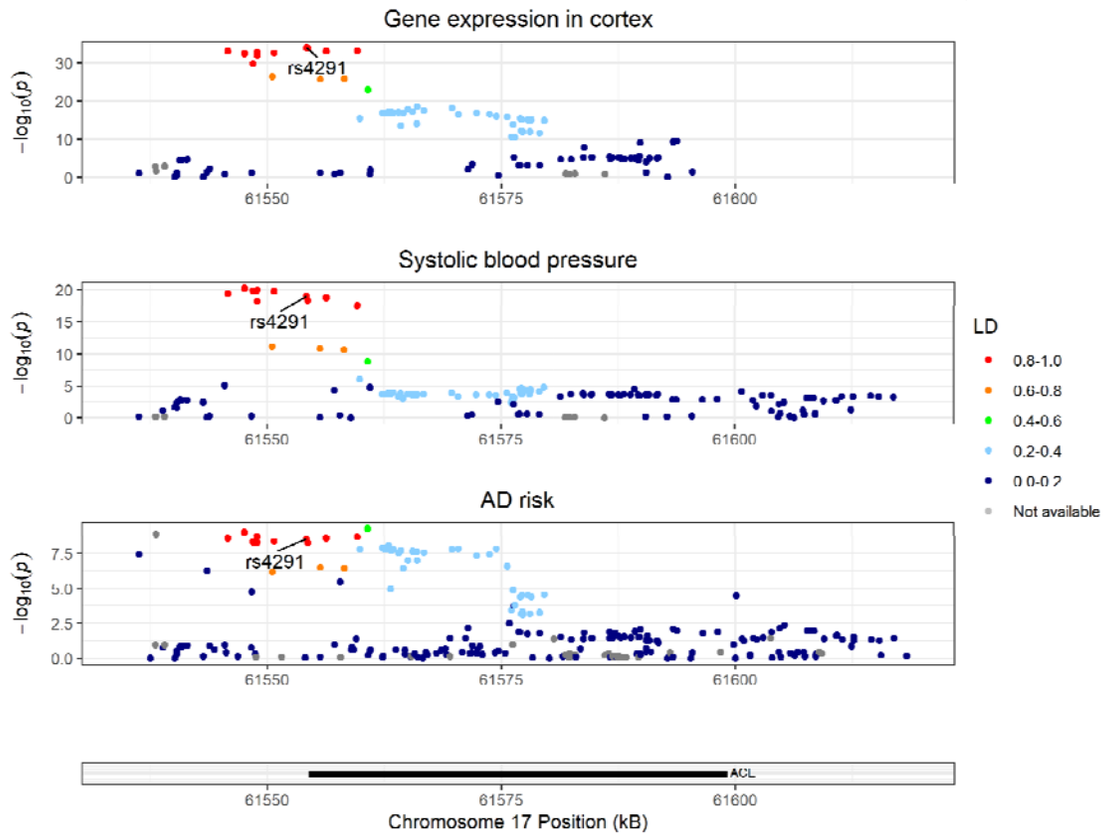


Figure 1: Colocalization plots depicting the association of single nucleotide polymorphisms (SNPs) with cortical ACE gene expression, systolic blood pressure and Alzheimer’s disease (AD) risk. The X axis shows position within the genome (build Hg19) and Y axis denotes the $-\log_{10}(p\text{-value})$ for the association. Colour denotes the linkage disequilibrium (LD) between different variants (see legend). The results support that rs4291 is the most likely candidate SNP underlying all these traits, therefore, representing a common causal SNP for these traits.

Associations between genetically proxied ACE inhibition and neurodegenerative traits

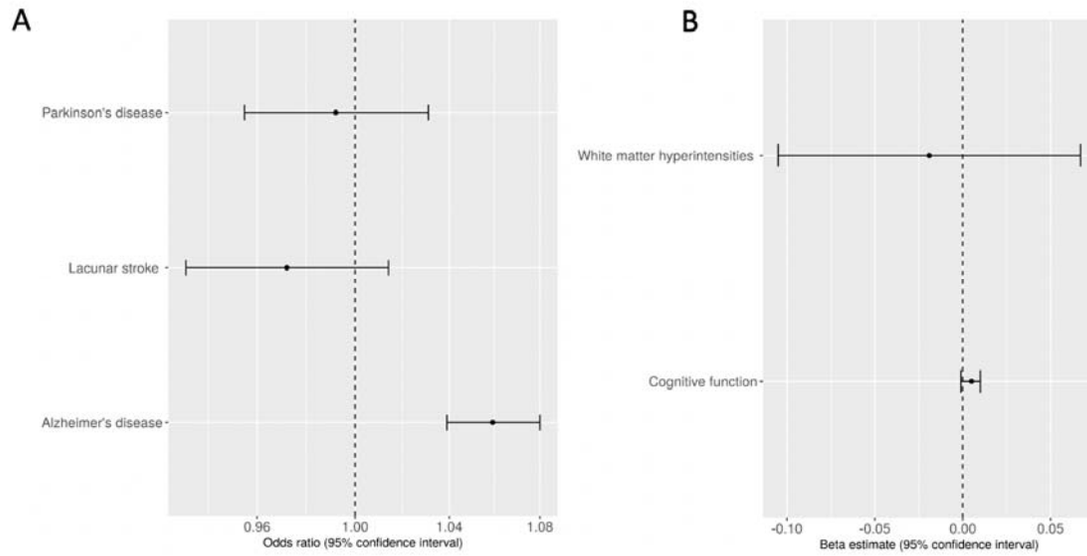


Figure 2: Forest plot showing associations between genetically proxied ACE inhibition (rs4291 effect allele A) with (A) disease outcomes and (B) continuous traits. In graph A, results reported as odds ratio per effect allele with 95% confidence interval. The x-axis is presented on the log10 scale. For graph B, results reported as beta estimate per effect allele with 95% confidence interval.