PLXNA4 is Associated with Alzheimer Disease and Modulates Tau Phosphorylation

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AUTHOR CONTRIBUTIONS G.J. designed and performed the genetic association analyses and co-wrote the paper. H.A. performed the tau phosphorylation experiments. E.Z. and C-D.C. performed the amyloid-β and APP experiments. E.D. performed the PLXNA4 isoform studies in brain. J.H.P, S.K., and J.W.K. designed primer sets and tested the efficacy in blood. T.F. and R.M. ascertained, phenotyped and genotyped the NIA-LOAD Study participants. V.H. obtained and characterized the brain specimens. R.K., M.A.P-V, J.L.H, and G.D.S provided Alzheimer Disease Genetics Consortium datasets and helped interpret results. K.L.L. consulted on the statistical genetic analyses and critically evaluated the manuscript. J.D.B. supervised the PLXNA4 genotyping and isoform studies in brain and critically evaluated the manuscript. T.I. designed and supervised the tau phosphorylation experiments and assisted in the preparation of the manuscript. C.R.A. designed the amyloid-β and APP experiments and assisted in the preparation of the manuscript. L.A.F. obtained funding for the study, coordinated all components of the study, evaluated and interpreted results, and co-wrote the paper.

CONFLICTS OF INTEREST Authors declare no conflict of interest.
Abstract

Objective—Much of the genetic basis for Alzheimer disease (AD) is unexplained. We sought to identify novel AD loci using a unique family-based approach that can detect robust associations with infrequent variants (minor allele frequency <0.10).

Methods—We conducted a genome-wide association study in the Framingham Heart Study (FHS) (discovery) and NIA-LOAD (replication) family-based cohorts using an approach that accounts for family structure and calculates a risk score for AD as the outcome. Links between the most promising gene candidate and AD pathogenesis were explored in silico as well as experimentally in cell-based models and in human brain.

Results—Genome-wide significant association was identified with a PLXNA4 SNP (rs277470) located in a region encoding the semaphorin-3A (SEMA3A) binding domain (meta-analysis p value [meta-P]=4.1×10^{-8}). A test for association with the entire region was also significant (meta-P=3.2×10^{-4}). Transfection of SH-SY5Y cells or primary rat neurons with full-length PLXNA4 (TS1) increased tau phosphorylation when stimulated by SEMA3A. The opposite effect was observed when transfected with shorter isoforms (TS2 and TS3). However, transfection of any isoform into HEK293 cells stably expressing APP did not result in differential effects on APP processing or Aβ production. Late-stage AD cases (n=9) compared to controls (n=5) had 1.9-fold increased expression of TS1 in cortical brain tissue (P=1.6×10^{-4}). Expression of TS1 was significantly correlated with the Clinical Dementia Rating score (ρ=0.75, P=2.2×10^{-4}), plaque density (ρ=0.56, P=0.01) and Braak stage (ρ=0.54, P=0.02).

Interpretation—Our results indicate that PLXNA4 has a role in AD pathogenesis through isoform-specific effects on tau phosphorylation.

INTRODUCTION

Alzheimer disease (AD) is the most frequent age-related dementia affecting 5.4 million Americans including 13% of people ages 65 and older and over 40% of people ages 85 and older. Genetic factors account for much of the risk for developing AD with heritability...
estimates between 60% and 80%. The apolipoprotein E (APOE) ε4 allele is a well-recognized major risk factor for late onset AD, increasing the odds of disease in a dose-dependent fashion. Common polymorphisms in 19 additional genes have been robustly established as risk factors for AD using large-scale genome-wide association studies (GWAS) and meta-analyses. These polymorphisms link to mechanisms of Aβ metabolism, lipid metabolism, inflammation, and axon guidance. However, the heritability of AD explained by APOE is less than 30% and by each of the novel GWAS loci is less than 1%, suggesting that less than 50% of the genetic contribution to AD is explained by known common polymorphisms. The remaining heritability may be due to additional common variants of weaker effect, rare variants, copy-number variants, insertion-deletion polymorphisms, and gene-gene and gene-environment interactions.

Here, we conducted a two-stage family-based AD GWAS using a novel method which incorporates the entire family structure and reduces diagnostic misclassification in the association test, and renders a result that is less prone to type I error, even for rare variants. We obtained strong evidence of association in the Framingham Heart Study (FHS) dataset with several SNPs in PLXNA4, a gene which had not been previously linked to AD. Several other PLXNA4 SNPs were highly significant in the National Institute on Aging-Late-Onset Alzheimer Disease (NIA-LOAD) study dataset. Subsequent in silico and molecular studies demonstrated isoform-specific effects of PLXNA4 on hyperphosphorylation of tau protein, a terminal step leading to breakdown of neuronal signaling and microtubule formation.

MATERIALS AND METHODS

Study Cohorts

**Framingham Heart Study (FHS) discovery cohort**—The FHS is a multigenerational study of health and disease in a prospectively followed community-based sample. Details on procedures for assessing dementia and determining AD status in this cohort are described elsewhere. We included only incident AD cases who had a magnetic resonance imaging (MRI) scan prior to disease onset. Clinical, demographic, genetic and pedigree information were obtained from dbGaP (http://www.ncbi.nlm.nih.gov/gap). Phenotypic and genome-wide association study (GWAS) data were available for 61 cases and 2,530 cognitively normal controls from 1,232 families. This sample contained 287 parent-offspring pairs, 1,215 sibpairs, 236 avuncular pairs, 714 cousin pairs, and 436 spouse pairs. The 61 AD cases are members of 57 families and three of these families contained distantly related affected individuals included in the analyses. Sixteen families did not have any genotyped AD cases but contained individuals with a history of dementia.

**National Institute on Aging – Late Onset Alzheimer’s Disease (NIA-LOAD) replication cohort**—The NIA-LOAD Study recruited families with two or more affected members available for genotyping. Phenotype and GWAS data for this cohort were obtained from dbGaP. The GWAS dataset included a total of 1,819 AD cases and 1,969 unaffected individuals from 2,265 families. The 1,819 AD cases are distributed among 988 families containing 1,010 concordantly affected, 878 discordantly affected, and 541
concordantly unaffected sibpairs. Within the GWAS dataset, there were 526 parent-offspring pairs, 2,429 sibpairs, 1,905 avuncular pairs, 1,533 cousin pairs, and spouse pairs.

Alzheimer’s Disease Genetics Consortium (ADGC) replication cohort—We obtained summarized results for SNPs located in the top-ranked locus from each of the individual ADGC datasets (excluding the NIA-LOAD dataset) which are described elsewhere. This sample included 18,901 Caucasians (9,966 cases and 8,935 controls), 4,896 African Americans (1,459 cases and 3,437 controls), and 1,845 Japanese (951 cases and 894 controls).

Genotyping, Quality Control, Population Substructure, and Imputation

FHS samples were genotyped at Affymetrix (Santa Clara, CA) using the Affymetrix GeneChip® Human Mapping 500K Array Set and 50K Human Gene Focused Panel®. NIA-LOAD samples were genotyped using Illumina 610 high-density SNP microarrays. APOE genotypes were obtained by restriction fragment length polymorphism analysis for the FHS cohort and by haplotype analysis of SNPs rs7412 and rs429358, which were genotyped at Prevention Genetics (http://www.preventiongenetics.com) for the NIA-LOAD cohort.

Quality control (QC) procedures and analysis of population substructure were performed as previously described. In the discovery genome-wide association (GWA) dataset, 341,492 SNPs genotyped in 2,591 subjects passed QC. Based on this number of SNPs, the threshold for genome-wide significance was $P=1.46\times10^{-7}$.

In candidate gene analyses in the FHS and NIA-LOAD datasets, we evaluated SNPs within 50 kb of the top-ranked gene from the discovery GWA study that were imputed previously using the HapMap 2 CEU reference panel and the MaCH program. SNPs with an imputation quality measure (RSQ)<0.80 were excluded.

Statistical Analysis

AD risk scores—We calculated AD risk scores using two different approaches. In the FHS dataset, liability scores are residuals from a logistic regression model for AD affection status adjusting for sex and censoring age (age at onset of AD or age at exam of controls) evaluated using R. We did not adjust for previously derived principal components (PCs) of population substructure because they were not significantly associated with the liability scores in the FHS dataset. In the NIA-LOAD dataset, we calculated liability scores (residuals) from a logistic regression model for AD affection status adjusting for sex, censoring age, and the first two genotype-covariance matrix PCs to account for population substructure. We also computed propensity scores in the NIA-LOAD dataset to account for the approximately 10-year lower mean age of this cohort compared to the FHS cohort and ascertainment of families with multiple affected members. This method was suggested previously for analysis of a binary trait with various ages at onset and exam among affected and unaffected subjects, respectively. Propensity scores were obtained for unaffected family members using the AGEON program in S.A.G.E. (v6.2) as the probability of getting disease conditioned on affection status and censoring age of their parents. Affection
status for AD cases (probability=1) was retained in the model. Propensity scores were further adjusted for censoring age, sex, and the first two PCs in a linear regression model, and the residuals used as the outcome variable. We were unable to obtain propensity scores in the FHS dataset because of incomplete parental information.

**SNP association test**—The quantitative risk scores (residuals from liability or propensity scores) were normalized by taking the inverse standard normal transformation of the empirical quantile that was obtained using the formula 
\[
\frac{r(y) - 1/3}{n+1/3}
\]
where \(r(y)\) is the rank of residuals, \(y\). This approach has been demonstrated to be valid for family-based association tests even for a binary trait with rare variants.\(^{23}\) Association of the normalized risk scores with each SNP was evaluated using the extension of the two-level Haseman-Elston regression method implemented in the RELPAL program in S.A.G.E. (v6.1).\(^{24,25}\) Models included a term for an inheritance vector of exhaustive relative pairs. Nominal \(P\) values for association were determined using first-level Wald tests. For top-ranked genes, we conducted association tests for genotype and well-imputed SNPs (\(RSQ \geq 0.8\)) in the FHS and the NIA-LOAD datasets using a risk score (i.e., liability score or propensity score) as a quantitative trait.

**Regional association test**—To allow for the possibility of allelic heterogeneity between the two family datasets, we performed association tests within each dataset for regions in the top ranked gene defined by the ligand-binding domain of the encoded protein using the Versatile Gene-Based Association Study (VEGAS) program which calculates \(P\) values based on 100,000 permutations.\(^{26}\)

**Meta-analysis**—Results from the FHS and NIA-LOAD datasets were combined by meta-analysis of \(Z\)-scores weighted by the number of subjects using METAL.\(^{27}\) In region-based meta-analysis, we assumed the same direction of effect on a region in both datasets and combined results using the \(Z\)-score approach. The summary results for the ADGC datasets (excluding the NIA-LOAD study) were meta-analyzed using the inverse variance method in METAL after applying a genomic control within each individual dataset.\(^{7}\)

**APP and Amyloid β (Aβ) Analyses**

**Cell maintenance and transfection**—HEK293 cells stably transfected with wild type APP751, were maintained as previously described.\(^{28}\) The day before transfection, cells were split into 6 well plates at a density of 1×10^5 cells per well. The cells were transfected with EV (empty vector) or PLXNA4 isoforms (TS1, TS2, or TS3) from OriGene using Attractene (Qiagen). Forty-eight hours post transfection, media were collected and cells were lysed as described.\(^{28}\)

**SDS-PAGE and western blotting**—Proteins were separated on 8% Tris-Glycine gels and transferred onto nitrocellulose (Millipore). Primary antibodies were mouse monoclonal 6E10 (Covance, 1:1000) against amino acids 1–17 of Aβ that also recognize full length APP and APPs\(\alpha\), anti-Myc mAb (Sigma, 1:1000) and mouse monoclonal beta-tubulin (Invitrogen, 1:10,000). Secondary antibody was peroxidase labeled goat anti-mouse IgG (KPL, 1:500). The Supersignal West Pico Chemiluminescent Substrate (Thermo Scientific)
was used for detection. Protein expression was analyzed by densitometry using ImageJ software. The expression of APPs\(\alpha\) was normalized to total APP.

**\(A\beta\) enzyme-linked immunosorbent (ELISA) assay**—Forty-eight hours after transfection of HEK293 cells (stably transfected with wtAPP751) with EV or PLXNA4 isoforms (TS1, TS2, or TS3), media were collected and centrifuged at 16,873g for 5min at 4°C to discard cell debris. ELISAs were carried out using the human A\(\beta\)40 and A\(\beta\)42 ELISA kits (Invitrogen) in accordance with manufacturer’s protocol with samples diluted 1:2 in diluent buffer and as previously described.\(^28\)

**Tau Analyses**

**Tau immunoblotting**—SH-SY5Y cells (2\(\times\)10\(^6\)) stably expressing mutant tau P301L (SH-SY5Y P301L) were transfected with 1 \(\mu\)g of various Plexin-A4-Myc or pcDNA3 vectors by lipofectamine 2000 (Invitrogen, Corp., Carlsbad, CA). After transfection, cells were treated with or without 3nM Semaphorin3A (Sema3A-FC, R&D Systems) for 1 hour. Twenty g of whole cell lysate were immunoblotted with phospho-tau (AT-8, 1:500), total-tau (Tau46, 1:500), \(\beta\)-actin (1:1000), or Myc (9E10, 1:1000) as previously described.\(^29\)

**Pull-down assay**—Plexin-A4-Myc transfected SH-SY5Y P301L cells were treated with 3nM 6xHis human Sema3A-FC in DMEM/F12/B-27 medium. After Sema3A-FC stimulation, the media were collected and the Sema3A-FC was precipitated with Protein A/G agarose, followed by immunoblotting of the samples for Myc (9E10, for detection of Plexin-A4 molecules) and 6x His (H-3, for detection of Sema3A, 1:1000, Santa Cruz Biotech).

**Primary neuron culture and tau immunofluorescence**—Rat primary hippocampal neurons were harvested from E18 embryonic brain, plated on poly-D-lysine coated coverslips, cultured for 14 days, and transfected with full-length or short isoforms of human Plexin-A4-Myc DNA plasmid vectors by lipofectamine 2000 (Invitrogen). Twenty-four hrs after transfection, cells were treated with or without 3 nM Semaphorin3A (Sema3A-FC) for 1-hr, fixed with 4% paraformaldehyde, and co-stained with T514 phosphorylated collapsing response mediator protein 2 (pT514 CRMP2, Ab62478, rabbit polyclonal 1:500, Abcam) and \(\beta\)3-tubulin (mouse monoclonal 1:500, Promega), or co-stained with AT8 (mouse monoclonal, 1:500) and anti-Myc (rabbit polyclonal, 1:200, Covance Inc), followed by incubation with anti-mouse IgG Alexa 568 conjugated goat secondary antibody (1:600, Invitrogen) and anti-rabbit IgG Alexa 488 conjugated goat secondary antibody (1:600, Invitrogen). Immunostained AT8 signals were digitally captured using an inverted fluorescence microscope (TE-2000U, Nikon Instruments, Melvin, NY) and quantified by ImageJ (NIH shareware program). Data were obtained from at least 100 cells in three independent experiments.

**Analyses of Gene Expression in Brain**

Gene expression experiments were conducted on brain tissue specimens obtained from 17 autopsied subjects including five controls (Braak stage 0), five early-stage AD cases (Braak stages 1–2), and nine late-stage AD cases (Braak stages 3–4). Ascertainment, cognitive
assessments, neuropathological assessments, and stratification of these subjects were previously described.\cite{30} Frozen post-mortem brain tissue specimens from Brodmann area 9 of subjects were obtained from the Mount Sinai/Bronx Veterans Administration (VA) Medical Center/Department of Psychiatry Brain Bank. Normal controls had no history of any psychiatric or neurological disorders and no discernible neuropathological lesions. The Institutional Review Boards of Pilgrim Psychiatric Center, the Icahn School of Medicine at Mount Sinai, and the Bronx VA Medical Center approved all assessment and post-mortem procedures.

Total RNA was isolated from 50 mg of tissue by using the Qiagen miRNeasy kit. Isolated RNA samples were treated with 40 units of DNase I (Ambion) to remove genomic DNA contamination. Template RNA quantity and quality, including degradation, were determined on an Agilent 2100 Bioanalyzer. Two samples with very low RNA integrity number were excluded from the study. After cDNA synthesis using Superscript III first-strand synthesis kit (Invitrogen, Carlsbad, CA), qPCR were run on an ABI 7900HT real-time machine in triplicate using Platinum Quantitative PCR SuperMix-UDG (Invitrogen, Carlsbad, CA) for primers designed with the Roche Universal Probe Library or SYBR GreenER qPCR supermix for regular primers for ABI PRISM (Invitrogen). Acquired data were loaded onto qBase ver.1.3.5 software for quality control and normalization to reference genes GAPDH and RPL13A.

Group differences between TS1 and TS3 or between controls and AD cases were evaluated using ANOVA and the appropriate T-test after checking for equality of variances. Pearson correlations were calculated using the SPSS program. Although post-mortem intervals (PMI) among AD cases and controls were comparable (P=0.450), we repeated the analyses including PMI as a covariate because differences in PMI can affect brain biochemistry and quality of RNA. Since results from models with and without PMI were similar, we reported only the results without adjustment for PMI.

**RESULTS**

**Genome-Wide Association**

The mean onset age of AD among the 61 incident cases in the FHS dataset was about 10 years older than that for the 1,819 AD cases in the NIA-LOAD dataset (Table 1). The frequency of APOE\(\varepsilon4\) carriers in affected subjects was approximately two times greater in the NIA-LOAD cohort compared to the FHS cohort. In addition, the proportion of APOE\(\varepsilon4\) carriers in controls was 15\% less in the FHS cohort. Among the 1,969 unaffected family members in the NIA-LOAD cohort, 439 had propensity scores of at least 80\% (i.e., risk to develop AD accounting for parental affection status and age at onset/exam) and 809 had propensity scores of zero indicating true controls.

Analysis of genotyped SNPs (n=341,492 post quality control) in the FHS dataset using the normalized liability scores for AD (Fig 1A) indicated little genomic inflation (\(\lambda=1.01\), Fig 1B) with strong evidence of association in three regions of the genome (Fig 1C). Genomewide significant associations were found with SNPs in ITIH3 (rs9311482; \(\beta=1.3,\) \(P=5\times10^{-9}\)), PLXNA4 (rs277484; \(\beta=1.1,\) \(P=9\times10^{-10}\)) and MYO18B (rs13057714; \(\beta=1.0,\) \(P=1.3\times10^{-7}\)).
SNPs located in *IGSF21* were suggestive at $P<10^{-6}$ (Supplementary Table 1). Each dose of the minor alleles for these SNPs increased AD liability by at least one rank unit (mean liability on the normalized scale). Based on an *ad hoc* power calculation using the sibling correlation of 0.2 derived from the data, the observed effect sizes and minor allele frequencies (MAF), and a genome-wide significance level of $1.46\times10^{-7}$, the FHS dataset had between 60% and 95% power to obtain these results even for infrequent SNPs ($0.01<\text{MAF}<0.05$) (Fig 2). Among the previously known genes for AD, one genotyped SNP in *BIN1* was moderately associated with the liability score ($rs10180840$, $P=1.4\times10^{-4}$). We attempted to discover additional association signals in *IGSF21, ITIH3, PLXNA4*, and *MYO18B* by evaluating 1,515 genotyped and accurately imputed SNPs (RSQ $\geq 0.8$) in the FHS dataset (Supplementary Table 2). *PLXNA4* SNPs $rs277470$, $rs277472$, $rs277476$, and $rs277484$ were significant ($P<10^{-9}$ for each, Fig 3A) and in complete linkage disequilibrium (LD) with each other (Fig 4). Among the top *PLXNA4* SNPs, only $rs277484$ was genotyped. No additional SNPs from the *IGSF21, ITIH3* or the *MYO18B* met the significance threshold of $1.4\times10^{-7}$. The dose of *APOE* $\varepsilon4$ alleles was not significantly associated with the AD liability score in FHS ($P=0.24$), perhaps because most of the incident cases in this cohort were observed at ages where the $\varepsilon4$ association with AD is relatively weak.\(^3\)

**Replication and Extended Analysis of *PLXNA4***

Next, we tested association of 746 genotyped and accurately imputed SNPs (RSQ $\geq 0.8$) from *PLXNA4* in the NIA-LOAD dataset (Fig 3B), but were unable to replicate any of the top-ranked SNPs obtained in the FHS sample (Supplementary Table 3). However, 16 of the 746 *PLXNA4* SNPs showed significant association at $P<10^{-3}$ in a model using the normalized liability score, and these association signals were improved using the normalized propensity score (Supplementary Table 3). We observed an association trend in the same effect direction with $rs277470$ in the NIA-LOAD using the propensity score ($p$ value: FHS=$2.1\times10^{-10}$, NIA-LOAD=$0.06$, meta-analysis=$4.1\times10^{-8}$) (Fig 3C). Each dose of the minor allele $C$ for $rs277470$ increased the liability rank by at least one unit in FHS dataset and the propensity rank by 13% in NIA-LOAD (Table 2). The most significant finding in NIA-LOAD (Fig 3B) was obtained with a genotyped SNP $rs12539196$ ($p$ value: FHS=$0.114$, NIA-LOAD=$3.7\times10^{-5}$, meta-analysis=$2.8\times10^{-5}$). The minor allele $C$ for $rs12539196$ decreased the liability rank by 0.09 in FHS and accounts for a 15% reduction in the propensity rank in NIA-LOAD (Table 2). The result for $rs12539196$ in the NIA-LOAD dataset was not meaningfully changed after adjustment for the number of *APOE* $\varepsilon4$ alleles ($P=2.7\times10^{-5}$) and remained significant after correction for multiple testing ($P=0.0036$).

Further scrutiny of the *PLXNA4* findings revealed that the most significant SNPs in each dataset are clustered in two distinct regions between recombination hot spots (Fig 3A to C), are separated by approximately 78,240 base pairs apart in intron 2 of the largest transcript (TS1), and flank an alternatively spliced exon present only in a much shorter transcript (TS3) (Fig 3D). Bioinformatic evaluation revealed that TS1 contains a transmembrane domain, whereas the shorter isoforms are predicted to be secreted (Fig 3E), suggesting that the longer and shorter isoforms may have distinct functional consequences related to AD. Based on this information, we performed region-based analyses including only SNPs located between 131,925,825 and 132,193,452 base pairs (including all of intron 2) encompassing...
the SEMA domain (Fig 3E), and confirmed significant association for the region (Table 2) in both datasets (p value: FHS=6.3×10^{-3}; NIA-LOAD=0.019; meta-analysis=3.2×10^{-4}).

Association with PLXNA4 was further examined using meta-analyzed results in the ADGC datasets. The most significant SNPs in each population under the additive model were rs10273901 in Caucasians (minor allele frequency [MAF]=0.42; meta-analysis p value [meta-P]=3.9×10^{-5}; odds ratio [OR]=0.85, 95% Confidence Interval [95% CI]: 0.79–0.92), rs75460865 in African Americans (MAF=0.04; meta-P=8.0×10^{-4}; OR=1.55, 95% CI: 1.20–2.01), and rs13232207 in Japanese (MAF=0.19; P=1.2×10^{-4}; OR=1.51, 95% CI: 1.22–1.86). The results for the Caucasian and Japanese groups remained significant after correcting for multiple testing (P=0.013 and P=0.037, respectively). Rs75460865 is located in the portion of the sequence which encodes the SEMA domain, but rs10273901 and rs13232207 are located in the PLXNA4 region that encodes the CYTO domain (Fig 3E). Top ranked SNPs in the FHS and NIA-LOAD datasets were not associated in the other ADGC datasets (P>0.1) likely because power to detect association with several SNPs (including rs277470) having low minor allele frequencies is weaker in the ADGC datasets compared to large extended family-based samples. Because the LD structure in this region is similar across populations (Fig 4), different association peaks among these groups is consistent with the existence of multiple distinct functionally-relevant AD-related alleles. In summary, the results in each of the ADGC ethnic samples support the association of AD with PLXNA4 SNPs.

**PLXNA4 Does Not Influence APP Processing or Aβ Production**

To determine whether PLXNA4 isoforms are differentially involved in APP processing and Aβ production, we transfected HEK293 cells stably expressing APP751 with empty vector control (EV) or with the full-length (TS1) or one of the shorter isoforms (TS2 and TS3) of PLXNA4-Myc and analyzed total APP in cell lysates, and APPs\(\alpha\), Aβ\(_{40}\), and Aβ\(_{42}\) in the medium. Neither over-expressing PLXNA4 isoforms affected APPs\(\alpha\) secretion (Fig 5A and B) or Aβ\(_{40}\) and Aβ\(_{42}\) production (Fig 5C). These results suggest that PLXNA4 is not involved in AD through the non-amyloidogenic or amyloidogenic processing of APP.

**PLXNA4 Isoforms Differentially Affect Tau Phosphorylation**

Involvement of plexin-A4 signaling in tau phosphorylation was examined by transfecting cDNAs for the full-length (TS1) or 3’ C-terminal truncated short isoforms (TS2 and TS3) of human PLXNA4-Myc into SH-SY5Y P301L cells (SH-SY5Y cells stably expressing the P301L tau mutant) and stimulated with or without 3 nM recombinant Semaphorin-3A (SEMA3A-FC). Immunoblotting with AT8 (anti-phospho-Tau at Ser\(^{202}/\text{Thr}^{205}\)) showed that tau phosphorylation was induced by SEMA3A stimulation, which was enhanced by over-expression of TS1 (Fig 6A). In contrast, over-expression of either TS2 or TS3 inhibited tau phosphorylation under stimulation by SEMA3A (Fig 6A). Since TS2 and TS3 are secretory molecules with the SEMA3A binding site in the extracellular domain, we assume that the inhibitory effect is mediated by the competitive binding of the short isoforms to SEMA3A. Pull-down assays confirmed that the short isoforms, but not full-length PLXNA4, specifically co-precipitated with SEMA3A in the media (Fig 3B). These data demonstrate that the short isoforms are secreted as expected and bind to SEMA3A thereby inhibiting
signaling. To confirm that SEMA3A/PLXNA4 signaling phosphorylates endogenous wild type tau protein, we repeated the experiment in rat primary hippocampal neurons. SEMA3A activated PLXN signaling leading to phosphorylation of endogenous CRMP2 at Thr 514 by glycogen synthase kinase-3β was confirmed in the primary cultured hippocampal neurons (Fig 6C). Stimulation of neurons with SEMA3A induced phosphorylation of endogenous CRMP2 at Thr 514 by glycogen synthase kinase-3β was confirmed in the primary cultured hippocampal neurons (Fig 6C). Stimulation of neurons with SEMA3A induced phosphorylation of endogenous tau as determined by AT8+ cells (Fig 6D, red staining in Myc− cells). Transient expression of Myc-tagged full-length PLXNA4 significantly elevated SEMA3A-induced endogenous tau phosphorylation (Myc+ AT8+ cells) as compared to untransfected neurons (Myc− AT8+ cells, p < 0.01), while expression of Myc-tagged shorter isoforms significantly reduced SEMA3A-induced tau phosphorylation in neurons (Fig 6D). This demonstrates that SEMA3A/PLXNA4 signaling phosphorylates endogenous tau in an isoform-specific manner.

Expression of PLXNA4 in Brain is Different in AD Cases and Controls

Expression of the TS1 and TS3 was quantified in brain tissue specimens from the middle frontal gyrus (Brodmann area 9) from 19 autopsied subjects including five controls, five early-stage AD cases, and nine late-stage AD cases (Supplementary Table 4) using the primer sets shown in Supplementary Table 5. Late-stage AD cases compared to controls had 1.9-fold increased expression of TS1 (P=6.0×10−4) and a more modestly increased expression of TS3 (P=0.021) and ratio of TS1/TS3 (P=0.066) (Fig 7). These patterns were similar in the comparison of all AD cases to controls (P-value: TS1=0.003, TS3=0.097, TS1/TS3=0.18) and are not age-related (Table 3). In the combined sample of AD cases and controls, TS1 level was significantly correlated with the clinical dementia rating score (r=0.75, P=2.2×10−4) and several measures of AD neuropathology (r≈0.5, P<0.05), but the correlations of TS3 level with these traits were much smaller (Table 3). These findings suggest that elevation in TS1 level increases risk for developing AD.

DISCUSSION

We identified significant association between AD risk and SNPs in PLXNA4 using a family-based approach. The top-ranked SNPs in the discovery and replication datasets are located in a single intron and surround an exon that is skipped in the processing of the full-length mRNA transcript. We also demonstrated that the full-length isoform (TS1), but not the shorter isoforms (TS2 and TS3) of PLXNA4 increased tau phosphorylation in SH-SY5Y cells stably expressing the P301L tau mutant and in primary rat neurons when stimulated by SEMA3A. Significantly higher levels of TS1 and TS3 in cortical brain tissue were observed in late-stage AD cases compared to controls. By comparison, transfection of either isoform into HEK293 cells stably expressing APP failed to show differential effects on APP processing or Aβ production. Taken together, our results indicate that PLXNA4-mediated tau phosphorylation is an independent upstream event leading to AD-related tangle formation in neurons.

PLXNA4 is a member of a family of receptors for transmembrane, secreted and GPI-anchored semaphorins in vertebrates and is a receptor for secreted semaphorin class 3 (SEMA3A) and class 6 (SEMA6) proteins which play an important role in semaphorin...
signaling and axon guidance. Accumulation of SEMA3A was previously detected in susceptible areas of the hippocampal neurons during AD progression and colocalized with phosphorylated tau. A phosphorylated CRMP2 protein, an intracellular signaling molecule for the semaphorin-plexin signaling pathway, has been observed in neurofibrillary tangles in brains of autopsied AD patients. The staining pattern of SEMA6, which is present in fibers and nerve terminals, is disrupted in brains of patients with AD. These reports and our study collectively indicate that disrupted semaphorin-plexin signaling is involved in AD pathogenesis, specifically through tau phosphorylation leading to tangle formation and neuronal death. Semaphorin-plexin signaling is known to regulate axon guidance in the development of sympathetic nervous system and cerebral cortex. Binding of SEMA3A to truncated PLXNA proteins was demonstrated to have a dominant negative effect on cortical growth cone collapse. Our data indicate that disruption of this signaling may also contribute to the acceleration of tau phosphorylation leading to neurofibrillary tangle formation. A recent whole exome sequence study identified a rare coding PLXNA4 variant in two distantly related individuals with familial Parkinson disease (PD). Notably, both PD subjects were cognitively impaired and there is a recognized form of PD with dementia. Further screening of all PLXNA4 exons in a large sample of unrelated PD cases and controls revealed a large number of very rare coding variants unique to cases or controls. Although the pathogenicity of these variants is yet unknown, this study provides additional evidence supporting PLXNA4 as a common biological pathway leading to AD and PD. Taken together, our findings point to a novel mechanism for AD-related tangle formation, implying that reduced expression of PLXNA4, and the TS1 isoform in particular, in brain is crucial to maintain healthy neurons.

There are three known alternatively spliced PLXNA4 transcripts. The full-length transcript (TS1) contains 31 exons and encodes an isoform with 1,894 residues. Two alternatively spliced transcripts each contain three exons yielding shorter isoforms of 522 residues (TS2) and 492 residues (TS3). The distinct PLXNA4 association peaks in the discovery and replication datasets flank an exon that is present only in TS3. A direct link of PLXNA4 expression to AD is supported by evidence in this report of increased expression of TS1 and TS3 isoforms in post-mortem neuronal tissue from AD cases compared to controls. Importantly, the findings that the relative increase of TS1 is much greater than TS3 in AD cases, and TS1 expression is significantly correlated with clinical and neuropathological severity measures of AD, are consistent with our observation of increased phosphorylation of tau by SEMA3A bound specifically to TS1.

Previous GWA studies involving the FHS dataset have successfully identified several AD genes including BIN1, MS4A4A/MS4A6A, EPHA7, ABCA7, CD33, CD2AP, HLA-DRB5–HLA-DRB1, PTK2B, SLCA24A4-RIN3, INPP5D, MEF2C, NME8, ZCWPW1, CELF1, FERMT2, and CASS4, but the portion of evidence for these associations attributable to the FHS dataset is very small and the robust associations with these loci are with common SNPs (MAF > 0.08) each exerting a small effect on risk (OR ≤ 1.2). We increased the potential for gene discovery in the FHS and NIA-LOAD datasets by applying an analytical method that leverages the family structure that is otherwise ignored when using Generalized Estimating Equations (GEE) models. Our approach is applicable to extended pedigrees and is robust for detecting association with infrequent SNPs (0.01 ≤ MAF<0.1) because the power
for this method is largely dependent on effect size rather than allele frequencies.\textsuperscript{11} Except for \textit{BIN1}, lack of significant associations with most of the previously known AD loci suggests that they do not contain large effect variants for AD risk in the range of infrequent MAFs. In addition, we addressed phenotype misclassification by including only incident cases with known onset ages in the FHS dataset and by using the propensity score approach in the NIA-LOAD dataset which has a skewed distribution of onset ages and is enriched with familial AD cases. Our method using propensity scores as a surrogate for AD susceptibility assigned probabilities greater than 50\% for future development of AD to 60\% of the unaffected subjects at last exam in the NIA-LOAD cohort. This indicates that analyses using affection status in families with multiple affected members have much less power to detect true associations without adjusting for misclassification of unaffected relatives.

Our study has several caveats. Power to detect association with AD risk is low in the FHS cohort because of the small number of incident cases. We overcame this limitation by using a continuous measure of AD liability instead of AD status as the outcome. In addition, the top-ranked PLXNA4 SNPs were not the same across datasets and thus the lack of an exact replication may indicate that the initial finding was a false positive. However, the inability to replicate at the SNP level can be explained by differences in allele frequencies across populations, and low power and reliability to detect association with infrequent SNPs in samples of unrelated cases and controls. Another likely contributor to the different association patterns is allelic heterogeneity, which we observed for \textit{SORL1}\textsuperscript{42} and has been reported among Caucasians for other common diseases.\textsuperscript{43, 44} Nevertheless, our PLXNA4 region-based test results were significant in both FHS and NIA-LOAD datasets. Moreover, the observed effects of PLXNA4 isoforms on AD-related processes in cultured neuronal cells and brain, which were identified through experiments conducted \textit{because} of the association findings, suggest that the significant results with individual PLXNA4 SNPs are not false positives. Another concern is that we did not have genotypes for the top ranked SNPs in a large sample of AD cases and controls with brain tissue available for study. As a result, we were unable to investigate allele specific effects on isoform expression levels. Moreover, demonstration of higher isoform levels in peripheral tissue of living AD cases compared to controls would lessen the concern that the differences observed in brain are not secondary to underlying pathology. In addition, \textit{in vitro} tau phosphorylation experiments do not indicate which tau kinases activate tau phosphorylation. Cycline-dependent kinase 5 and glycogen synthase kinase-3\(\beta\) are known to be activated by SEMA3A by phosphorylating CRMP2 to mediate growth cone collapse.\textsuperscript{45, 46} It is most likely that PLXNA4 induces tau phosphorylation via activation of these two established kinases. Also, because these results are based on the transient expression of PLXNA4 molecules, it is crucial to evaluate the influence of PLXNA4 on tau phosphorylation in more physiological conditions. Isoform-specific targeting of \textit{PLXNA4} in mice is a potential future study \textit{in vivo}.

In summary, our novel genetic association findings and results of molecular and cell biology experiments in cell lines, rat neurons, and human brain demonstrate that \textit{PLXNA4} is involved in AD pathogenesis. Evidence supporting transcriptional regulation of \textit{PLXNA4} isoforms that have differential effect on tau phosphorylation and, hence, tangle formation suggests the potential for a new drug target. These findings warrant follow-up experiments to delineate the upstream factors governing PLXNA4 binding to CRMP2, ascertain the
causative genetic variants at the two association peaks and determine how they modulate expression of the individual isoforms, demonstrate in brain tissue that the genetic variation in PLXNA4 is associated with AD risk or progression, and evaluate the utility of PLXNA4 as a potential biomarker for AD. Further studies of PLXNA4 may also provide mechanistic evidence linking AD to PD with comorbid dementia.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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REFERENCES


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Fig 1. Genome-wide association analysis in the Framingham Heart Study

(A) Ranked risk score distribution. Liability scores after adjusting for age and sex in a logistic regression model were rank-transformed and analyzed as a quantitative trait. Areas with red indicate distribution of risk (liability) rank of AD cases based on age at onset. Black arrow for age at exam from controls and red arrow for age at onset from AD cases indicate ranks of risk scores. The curve shows younger unaffected subjects have higher risk to develop AD than older unaffected subjects, and AD cases with earlier onset have greater liability than cases with older onset. (B) Quantile-quantile plot. Observed P values (Y-axis) were plotted against expected P values (X-axis). Black dots represent P values for all genotyped SNPs. (C) Manhattan plot. Association results for genotyped SNPs. P-values are expressed as $-\log_{10}(P)$ (y-axis) for every tested SNP ordered by chromosomal location (x-axis). Genome-wide significance level is shown as a dotted line at $P=1.4\times10^{-7}$. 
Fig 2. Power to detect genome-wide significant association with AD in the Framingham Heart Study

Effect sizes (i.e., beta estimates in the regression model) to obtain 80% power for infrequent SNPs (0.01 ≤ minor allele frequency [MAF] ≤ 0.1) under the additive model were estimated using a sibling correlation of 0.199 in 2,779 relative pairs at the genome-wide significance level. Effect size of the SNP on AD risk (Y-axis) according to MAF (X-axis) was computed for rank-transformed liability scores. The power estimates are conservative because they account for only sib-pair relationships in the pedigree.
Fig 3. Genetic findings in the PLXNA4 region

Regional association plots of genotyped and imputed SNPs from the FHS (A) and NIA-LOAD (B) datasets, and in meta-analysis (C). Most significant SNPs in the FHS (rs277470) and NIA-LOAD (rs12539196) datasets are indicated by purple diamonds. P-values are expressed as \(-\log_{10}(P)\) (y-axis) for every tested SNP ordered by chromosomal location (x-axis). Estimates of linkage disequilibrium \((r^2)\) of SNPs in this region with the top SNP computed using 1000 Genomes (hg19/Nov2010EUR) are shown as orange circles for \(r^2 \geq 0.8\), yellow circles for \(0.5 \leq r^2 < 0.8\), light blue circles for \(0.2 \leq r^2 < 0.5\), and blue circles for \(r^2 < 0.2\). Recombination rates (scale on right axis) are plotted with a solid gray line.

Genomic structure of PLXNA4 was determined using the NCBI database (Build 37.1). (D) Relative position of the most significantly associated SNPs in FHS and NIA-LOAD datasets in the three validated transcripts (TS1, TS2, and TS3). Exons are denoted with horizontal bars. (E) Diagrams of functional domains encoded by amino acids from the full-length (TS1) and the shorter (TS2 and TS3) transcripts. SEMA: sema_plexinA1 interacting module (exon 1–4 in TS1 and exon 1–3 in TS2 and TS3); PSI: plexin repeat (exon 4–11 in TS1); IPT: three repeats of the binding domains of plexins and cell surface receptors (exon 11–17 in TS1); PCSR: binding domain of plexins and cell surface receptors (PCSR) and related proteins (exon 17–19 in TS1); TM: transmembrane region (exon 19 in TS1); CYTO: cytoplasmic domain (exon 20–31 in TS1).
Fig 4. Linkage disequilibrium (LD, D') of top ranked SNPs

LD was calculated in 1000 Genomes data from (A) Caucasians (CEU), (B) African Americans (AA), and (C) Asians (ASN). Top-ranked SNPs from PLXNA4 from each dataset are shown, namely rs277470, rs277472, and rs277484 in FHS; rs12539196 in NIA-LOAD, rs10273901 in ADGC Caucasians (ADGC-EA), rs75460865 in ADGC African Americans (ADGC-AA), and rs13232207 in ADGC Japanese (ADGC-JPN). The top SNP in the ADGC-AA dataset was monomorphic in both CEU and ASN populations. Five top-ranked SNPs (rs10273901, rs75460865, rs277470, rs277472, and rs277484) from the ADGC-EA and ADGC-AA samples were monomorphic in the ASN population. The top ranked SNPs are located in the SEMA domain, except rs10273901 and rs13232207 which are located in the cytoplasmic domain.
Fig 5. Effect of expression of the full-length PLXNA4 isoforms (TS1, TS2 and TS3) on APP processing

HEK293 cells stably overexpressing APP were transiently transfected with empty vector control (EV) or with the full-length (TS1) or one of the shorter isoforms (TS2 and TS3) of PLXNA4-Myc. 48 hrs after transfection, the conditioned medium (medium) and the cell lysates (lysate) were collected and analyzed by SDS-PAGE and western blotting using mAb 6E10 for total APP and APP$_{\alpha}$, Myc mAb for all 3 isoforms of PLXNA4, and tubulin as control (A). The endogenous myc can be seen in all lanes at ~55 kDa but more clearly in lanes 7–9 where only empty vector was transfected but not PLXNA4-myc. Also note that TS1 migrates at 212kDa, TS2 migrates at 58kDa and TS3 migrates at 55kDa. (B) Densitometric analysis of the expression of APP$_{\alpha}$ normalized to total APP. Error bars indicate standard deviation. (C) ELISA analysis of A$\beta_{40}$ and A$\beta_{42}$ released to the medium. ELISAs were carried out using the human A$\beta_{40}$ and A$\beta_{42}$ ELISA kits (Invitrogen) in accordance with manufacturer’s protocol. Error bars indicate standard deviation. Representative results of three independent experiments are shown in each panel.
Fig 6. Effect of PLXNA4 isoforms on tau phosphorylation
(A) SH-SY5Y P301L cells were transfected with the full-length (TS1) or one of the shorter isoforms (TS2 and TS3) of PLXNA4-Myc or empty vectors (pcDNA3.1) with or without 3nM SEMA3A stimulation for 1hr. Whole cell lysates were blotted with AT8, total tau, actin and Myc. Results for the TS2 and TS3 isoforms were similar but only those for TS3 are shown. (B) 6x His-tagged SEMA3A-Fc was precipitated from media by Protein A/G agarose, and the precipitates were immunoblotted with antibodies to Myc (detecting PLXNA4 isoforms) and 6x His (detecting SEMA3A-FC). (C) E18 rat primary hippocampal neurons were cultured for 14 days and stimulated with 3nM SEMA3A for 1 hr, and co-stained with anti-β3-tubulin (B3T) mouse monoclonal (1:500, green) and anti-pT514 CRMP2 rabbit polyclonal (1:500, red). (D) Rat primary hippocampal neurons were transfected with the full-length or short isoforms of Plexin-A4-Myc. After transfection, cells were treated with or without 3 nM Sema3A for 1-hr, and co-stained with AT8 (1:500) and anti-Myc (rabbit polyclonal, 1:200, Covance Inc). AT8 signals were digitally captured and quantified as previously reported (Uchida et al., 2005). Cells are immunostained with anti-Myc (green) and AT8 (red). Scale bar represents 10 μm. * P<0.05, ** P<0.01, and *** P<0.001, as determined by ANOVA and Tukey post hoc.
Fig 7. Expression of *PLXNA4* isoforms in post-mortem brains

Expression of *PLXNA4* isoforms from Broadmann area 9 of frozen post-mortem brain tissue specimens were compared in controls and late-stage AD cases using primer sets to detect full-length (TS1) and short (TS3) isoforms. RNA expression level (Y-axis) shown is the normalized value. P-values were determined by T-test accounting for unequal variances.
### Table 1

Sample characteristics

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Family Set</th>
<th>ADGC Set</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>FHS</td>
<td>NIA-LOAD</td>
</tr>
<tr>
<td>Total, N</td>
<td>2876</td>
<td>3828</td>
</tr>
<tr>
<td>AD Cases, N</td>
<td>61</td>
<td>2530</td>
</tr>
<tr>
<td>Female %</td>
<td>54</td>
<td>63</td>
</tr>
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<td>Age at onset (AD cases), mean (SD)</td>
<td>82.8 (9.2)</td>
<td>73.3 (7.0)</td>
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<tr>
<td>Age at last exam (AD controls), mean (SD)</td>
<td>68.1 (10.6)</td>
<td>75.4 (11.6)</td>
</tr>
<tr>
<td>% of APOE ε4-carriers in cases</td>
<td>42.6</td>
<td>75.0</td>
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<tr>
<td>% APOE ε4-carriers in controls</td>
<td>21.6</td>
<td>36.6</td>
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</table>

FHS=Framingham Heart Study; NIA-LOAD=National Institute on Aging-Late Onset Alzheimer’s Disease. ADGC replication set consisted of previously reported unrelated datasets from 14 Caucasian studies (excluding NIA-LOAD), 7 African American datasets (excluding NIA-LOAD), and one Japanese dataset.
Table 2

Top-ranked SNP-based association results for *PLXNA4* in the FHS and NIA-LOAD datasets

<table>
<thead>
<tr>
<th>SNP or Region</th>
<th>BP</th>
<th>RA</th>
<th>FHS</th>
<th>NIA-LOAD</th>
<th>Meta-Analysis</th>
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</thead>
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<tr>
<td></td>
<td></td>
<td></td>
<td>RAF  β  P-val</td>
<td>RAF  β  P-val</td>
<td>Z  P-val</td>
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<tr>
<td>rs277470</td>
<td>132,110,922</td>
<td>C</td>
<td>0.01 1.15 2.1×10^{-10}</td>
<td>0.01 0.13 0.0616</td>
<td>5.49 4.1×10^{-8}</td>
</tr>
<tr>
<td>rs12539196</td>
<td>132,037,683</td>
<td>A</td>
<td>0.84 0.09 0.114</td>
<td>0.85 0.15 3.7×10^{-5}</td>
<td>4.19 2.8×10^{-5}</td>
</tr>
<tr>
<td>SEMA Region*</td>
<td>131,925,825 – 132,193,452</td>
<td>nSNP=246 6.3×10^{-3}</td>
<td>nSNP=240 0.019</td>
<td>3.60 3.2×10^{-4}</td>
<td></td>
</tr>
</tbody>
</table>

FHS=Framingham Heart Study; NIA-LOAD=National Institute on Aging Late Onset Alzheimer’s Disease; N=sample size; BP=map position in base pairs from CRCh37/hg19; RA=reference allele; RAF=reference allele frequency; β=estimate of effect size; P-val=p value; Z=Z-score in meta-analysis; nSNP, the number of SNPs in a region based test.

*Region-based test was conducted using all SNPs from 131,925,825 to 132,193,452 base pairs flanking exons 1 to 3 from three transcript isoforms on chromosome 7.
Table 3
Correlation of PLXNA4 isoform levels with clinical and neuropathological phenotypes in brain specimens from subjects with AD and controls.

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Correlation (P value)</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>TS1</td>
</tr>
<tr>
<td>Age</td>
<td>0.27 (0.27)</td>
</tr>
<tr>
<td>PMI</td>
<td>-0.01 (0.97)</td>
</tr>
<tr>
<td>RIN</td>
<td>-0.09 (0.70)</td>
</tr>
<tr>
<td>CDR</td>
<td>0.75 (2.2x10^-4)</td>
</tr>
<tr>
<td>Plaque BM9</td>
<td>0.47 (0.045)</td>
</tr>
<tr>
<td>Plaque Meanc</td>
<td>0.56 (0.012)</td>
</tr>
<tr>
<td>Braak Stage</td>
<td>0.54 (0.017)</td>
</tr>
</tbody>
</table>

PMI=postmortem interval; RIN=RNA integrity number; CDR=clinical dementia rating; Plaque BM9=neuritic plaque density Brodmann area 9; Plaque Mean=mean of plaque densities measured in five cortical regions including Brodmann area 9 (middle frontal gyrus), Brodmann area 45/47 (orbital frontal gyrus), Brodmann area 21/22 (superior temporal gyrus), Brodmann area 39 (inferior parietal cortex), and Brodmann area 17 (calcarine cortex).