Segregation of a rare \textit{TTC3} variant in an extended family with late-onset Alzheimer disease

\textbf{ABSTRACT}

\textbf{Objective:} The genetic risk architecture of Alzheimer disease (AD) is complex with single pathogenic mutations leading to early-onset AD, while both rare and common genetic susceptibility variants contribute to the more widespread late-onset AD (LOAD); we sought to discover novel genes contributing to LOAD risk.

\textbf{Methods:} Whole-exome sequencing and genome-wide genotyping were performed on 11 affected individuals in an extended family with an apparent autosomal dominant pattern of LOAD. Variants of interest were then evaluated in a large cohort of LOAD cases and aged controls.

\textbf{Results:} We detected a single rare, nonsynonymous variant shared in all 11 LOAD individuals, a missense change in the tetratricopeptide repeat domain 3 (\textit{TTC3}) gene. The missense variant, rs377155188 (p.S1038C), is predicted to be damaging. Affecteds-only multipoint linkage analysis demonstrated that this region of \textit{TTC3} has a LOD score of 2.66 in this family.

\textbf{Conclusion:} The \textit{TTC3} p.S1038C substitution may represent a segregating, rare LOAD risk variant. Previous studies have shown that \textit{TTC3} expression is consistently reduced in LOAD patients and negatively correlated with AD neuropathology and that \textit{TTC3} is a regulator of Akt signaling, a key pathway disrupted in LOAD. This study demonstrates how utilizing whole-exome sequencing in a large, multigenerational family with a high incidence of LOAD could reveal a novel candidate gene.

\textbf{GLOSSARY}

\textit{AD} = Alzheimer disease; \textit{ADGC} = Alzheimer’s Disease Genetics Consortium; \textit{APOE} = apolipoprotein \textit{E}; \textit{APP} = amyloid precursor protein; \textit{ESP} = Exome Sequencing Project; \textit{FTD} = frontotemporal dementia; \textit{LOAD} = late-onset AD; \textit{MAF} = minor allele frequency; \textit{PSEN} = presenilin; \textit{SNV} = single-nucleotide variant; \textit{TTC3} = tetratricopeptide repeat domain 3; \textit{WES} = whole-exome sequencing.

Alzheimer disease (AD) is the most common form of dementia in the elderly, accounting for up to 75% of all dementia patients.\textsuperscript{1} Mutations in the amyloid precursor protein (\textit{APP}), presenilin 1 (\textit{PSEN1}), and presenilin 2 (\textit{PSEN2}) genes have been shown to cause autosomal dominant, early-onset AD.\textsuperscript{2–4} Late-onset AD (LOAD) is etiologically more complex and presents with intermediate to high heritability (58%–79%).\textsuperscript{5} The apolipoprotein \textit{E} (\textit{APOE}) gene contributes significantly to LOAD risk; the \textit{APOE} \textit{e4} allele confers an increased risk to LOAD, while the rare \textit{APOE} \textit{e2} allele is protective.\textsuperscript{6–9} In addition to the \textit{APOE} gene, genome-wide association studies identified 19 further loci that confer low individual, but reproducible risk to LOAD.\textsuperscript{10} Moreover, rare variants in \textit{TREM2} have been shown to contribute to LOAD risk.\textsuperscript{11} It seems likely that additional, yet unknown, rare variants with intermediate to large effects may

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 contribute to the complex genetic architecture of LOAD. We identified a large, multiplex LOAD family with an autosomal dominant pattern of inheritance and negative for pathogenic variants in known AD genes. We performed whole-exome sequencing (WES) in 11 affected family members and prioritized rare variants that segregated with disease. A single alteration in tetratricopeptide repeat domain 3 (TTC3) was identified in all 11 LOAD individuals. Our study shows that WES in a single extended family with a high LOAD burden can lead to the identification of rare genetic variants that may contribute to risk for AD.

METHODS Standard protocol approvals, registrations, and patient consents. All individuals ascertained in this study provided written informed consent prior to their participation. If the study participant was not competent to provide consent, then written consent was obtained from the immediate next of kin or his or her legal representative. All participants were ascertained using the protocol approved by the appropriate institutional review board. Oversight of the study falls under the University of Miami (IRB #20070307).

Collection and characterization of an extended LOAD family. We identified a large family (no. 1229) that was consistent with an autosomal dominant inheritance pattern of LOAD. Family 1229 has 15 affected individuals, 3 of whom have a neuropathologically confirmed diagnosis of LOAD. All affected individuals meet the standard National Institute of Neurological and Communicative Disorders and Stroke and Alzheimer Disease and Related Disorders Association (NINCDS-ADRDA) criteria for LOAD with a disease onset of 65 years of age or older.12 Besides direct neurologic examinations, a battery of neuropsychiatric testing was performed to monitor preclinical symptoms of LOAD and comorbidities (Parkinsonism, depression) that are known to influence the course of preclinical AD. The family was evaluated to determine APOE ε genotypes and screened for high-risk variants in APP, PSEN1, and PSEN2.

Genome-wide genotyping and linkage analysis. Genotype data were obtained for 9 LOAD-affected individuals of family 1229 using the Human OmniExpress BeadChip array (Illumina, San Diego, CA). Genotype quality control comprised the exclusion of single-nucleotide variants (SNVs) with call rates below 98%. The B allele frequency plots and the log R ratio plots were used to search for potential copy number variants surrounding our top variant. These data were analyzed with MERLIN to determine an affecteds-only multipoint linkage LOD score with LOAD in family 1229 assuming an autosomal dominant model.13 Two-point age-dependent penetrance linkage was also performed for the variant of interest using an age curve as previously described.14

Whole-exome sequencing. WES was performed in all 11 affected individuals of family 1229. WES used 3 µg of DNA from each sample. Exome capture and sequence library construction were performed on a Sciclone G3 NGS Workstation (Caliper Life Sciences, Hopkinton, MA) using the SureSelect Human All Exon 50 Mb Kit (Agilent Technologies, Santa Clara, CA) and the Paired-End Multiplexed Sequencing library kit (Illumina) for sequence library preparation. Two exome sample libraries were sequenced per lane on a HiSeq 2000 Sequencing System (Illumina) in paired-end 2×100 base pair runs. In addition, WES of 373 non-AD samples of European ancestry were also run and analyzed at the John P. Hussman Institute for Human Genomics in order to get minor allele frequency (MAF) estimates and control for WES artifacts.

SNV and insertion-deletion calling and annotation. Sequencing data were processed using the Illumina RTA base calling pipeline v1.8. Alignment to the human reference genome (hg19) was executed with the Burrows-Wheeler Aligner and variant calling performed with the Genome Analysis Toolkit (GATK).15–16 GATK parameters for variant quality control included duplicate sequence read removal, base quality score recalibration (QUAL >50), and variant quality score recalibration (VQSLOD >0).17 SNVs passing quality measurements were annotated with SeattleSeq (http://snp.gs.washington.edu/SeattleSeqAnnotation137/). We define rare SNVs and insertions-deletions (indels) as those with a MAF <1% in dbSNP139 (http://www.ncbi.nlm.nih.gov/snp/), 1000 Genomes Project populations, the 5,379 European and African-American samples from the NHLBI GO Exome Sequencing Project (ESP), Seattle, WA (http://evs.gs.washington.edu/EVS/), and 373 Caucasian exomes processed at the John P. Hussman Institute for Human Genomics.18,19 Variants were evaluated for phylogenetic conservation in mammals with the Genomic Evolutionary Rate Profiling score and PhastCons.20–21 A Genomic Evolutionary Rate Profiling score >3 and a PhastCons score >0.7 were considered phylogenetically conserved. A variant’s predicted deleteriousness to protein function was assessed by the Likelihood Ratio Test, the Mutation Assessor, Mutation Taster, PolyPhen-2 (HumDiv), and PROVEAN/SIFT programs using the default score cutoffs for deleteriousness.22–26

SNV validation by Sanger sequencing. Ten rare SNVs identified by WES that passed quality control and were shared in >80% (≥9/11) of affected individuals of family 1229 were evaluated by Sanger sequencing in the 11 affected and 5 unaffected members of this family.

Evaluating the TTC3 variant in a LOAD cohort. The TTC3 variant was evaluated using a custom TaqMan genotyping assay (Life Technologies, Carlsbad, CA). A large cohort of 6,669 LOAD cases and 5,585 controls (≥60 years) of European ancestry from the Alzheimer’s Disease Genetics Consortium (ADGC) cohort were used. Power calculations were performed with the Genetic Power Calculator.27

RESULTS Clinical and neuropathologic characterization. DNA samples were collected from family 1229 from 11 affected and 5 unaffected members of at least 65 years old (figure 1, table e-1 at Neurology.org/ng). Clinically, each of the 11 affected individuals presented with late-onset dementia (range: 70–85 years old). The initial course of disease followed a typical AD pattern, with early forgetfulness and progressive worsening in areas of cognitive function such as short-term memory, orientation, executive function as reflected in activities of daily living, and language retrieval (word finding).

Patients were assessed for features suggestive of alternate demyelinating pathologies, such as dementia.
with Lewy bodies, frontotemporal dementia, and vascular dementia. Abnormal features suggestive of a frontotemporal etiology such as frontal release signs, early changes in personality, behavior, or language changes were not present in any family members. A positive snout reflex was noted approximately 9 years into the disease course in one individual (0122), but no other individual exhibited frontal release signs or symptoms. The late appearance of a single frontal release sign is not suggestive of a diagnosis of frontotemporal dementia. A second individual (9006) exhibited mild parkinsonian features (masked facies, hunching, micrographia, poor balance) when examined at age 84, ten years into the disease course. The later onset of comorbid parkinsonian features is a known entity in AD.

Four affected individuals underwent autopsy with subsequent neuropathologic results (table e-1). All individuals demonstrated classic Braak and Braak pathology, ranging in stage from III to V.

Complete segregation of the rare TTC3 p.S1038C substitution with LOAD. WES in 11 LOAD members of family 1229 resulted in an average coverage of 83% (±0.04% SD) of the human exome sequence at a minimal depth of 20 reads per base pair. On average, 11,227 (±305 SD) exonic SNVs were called per exome, for a total of 32,784 unique SNVs across all 11 affected family members. The majority of disease-causing variants identified to date are nonsynonymous, protein-altering SNVs including nonsense, missense, and splice site variants as well as small insertion-deletion polymorphisms (insertions/deletions). Of all exonic variants detected in family 1229, about half (16,363 SNVs) were found to be nonsynonymous.

Family 1229 is characterized by a suggestive autosomal dominant pattern of LOAD inheritance (figure 1). We therefore hypothesized that a single (or a few) highly penetrant disease variant, with considerable individual risk, could explain the high burden of LOAD observed in this family. Statistical power estimates for the detection of LOAD variants suggest that published genome-wide association studies would likely have identified loci that harbor variants with aMAF >1% in the general population. Thus, we further filtered for rare variants (MAF <1%), which reduced the candidate variants to 2,897 SNVs. Among these rare and nonsynonymous variants, only a single variant, rs377155188, was identified in all 11 affected family members; this variant is in the TTC3 gene and encodes for a serine to cysteine substitution at amino acid position 1038 (figure 1; table 1).

We evaluated the remaining rare, nonsynonymous variants. None were shared in 10 of 11 affected family members, but 9 variants were shared in 9 affected family members (table 1). All 10 variants in at least 9 LOAD family members were confirmed by traditional Sanger sequencing (table 1).

The TTC3 variant, rs377155188, is extremely rare in European Americans (MAF = 0.00012) and absent in African Americans according to the NHLBI ESP. The p.S1038C substitution in the TTC3 protein is predicted to be damaging by 5 distinct algorithms. Furthermore, it is located in a highly conserved site.
The *TTC3* heterozygous genotypes identified through WES were validated by Sanger sequencing in all 11 affected individuals. Wild-type genotypes were obtained for 4 out of 5 unaffected family members ≥65 years old (figure 1). Individual no. 9018 was the only affected family member confirmed to harbor a heterozygous carrier of *TTC3* rs377155188. However, this individual did not receive a formal cognitive examination as did the 3 other unaffected family members. This individual’s negative affection status of LOAD has been based on family history data after the individual’s death at age 76. This person’s age at death is only 1 year above the average age of LOAD onset in this family (table e-1); thus, given the absence of a formal examination, this carrier may still have been symptomatic at age 76.

We then evaluated the prevalence of rs377155188 by genotyping 12,254 unrelated individuals (6,669 LOAD cases and 5,585 controls) of European ancestry from the ADGC. Over all individuals, rs377155188 showed a low frequency (MAF = 0.020%), as has been reported by the ESP for Americans of European descent (MAF = 0.012%).

We observed 5 heterozygous individuals in total. Four were LOAD cases, all of whom carried the *APOE* ε3/ε4 genotype. Only a single cognitively normal control carried the *TTC3* variant (0.018%). This individual was last examined at age 84 and also carried a protective *APOE* ε2/ε3 genotype. These genotypes resulted in an allelic association odds ratio of 3.35, which did not reach statistical significance (Fisher exact test, *p* = 0.248). Power calculations show that we are underpowered to detect a significant effect in our cohort (49.6% power to achieve significance at *α* = 0.05; assuming a disease frequency of 0.10), given the extremely rare allele frequency of the rs377155188 variant in our cohort and the general population.

**DISCUSSION** We performed WES in an extended, multigenerational family with an autosomal dominant inheritance pattern of LOAD. Disease segregation analyses identified the *TTC3* p.S1038C substitution as the only rare, nonsynonymous variant completely shared among all LOAD family members. It reached a suggestive multipoint linkage LOD score of 2.66. Moreover, it was absent in 3 aged family members with confirmed normal cognition (figure 1). A fourth affected family member carried the variant in *TTC3*, but died at age 76, very close to the average age at onset for the 11 AD relatives with the variant, 75.7 years old. Therefore, it is possible that if this individual had lived longer, he may have developed AD. Thus, the *TTC3* variant may, like *APOE*, contribute to LOAD risk; however, it is unlike *APP*, *PSEN1*, and *PSEN2* and may not be sufficient to initiate AD pathogenesis.

LOAD is a common, complex, and age-related disease. Additional factors, such as the aggregation of several genetic risk variants, might contribute to LOAD. Individual 122 presented with the *TTC3* risk allele and an *APOE* ε3/ε3 genotype, demonstrating that the presence of an *APOE* ε4 is not required to confer AD risk in this family (figure 1; table e-1). In addition, the family has 9 ε3/ε4 AD carriers and a single ε4/4 affected individual. However, the high abundance of *APOE* ε3/ε4 genotypes in family 1229 may not be sufficient to explain the apparent autosomal dominant inheritance pattern of LOAD. The average age of LOAD onset among affected family members was 75.7 years. A study representative of the US population determined the prevalence of LOAD at ≥70 years of age to be 9.7% (95% confidence interval: 7.6%–11.9%). The LOAD risk of *APOE* ε3/ε4 carriers is about 3-fold higher than that found in ε3/3 carriers, while ε4/4 carriers have a 15-fold higher risk of AD. We would expect a few phenocopies attributable to the *APOE* ε4 allele in family 1229 by assuming that the single ε4/4 carrier and

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**Table 1** Top rare LOAD candidate variants identified in family 1229

<table>
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<th>Gene</th>
<th>Genomic location [hg38]</th>
<th>dbSNP</th>
<th>Nucleotide</th>
<th>Amino acid change</th>
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<td>Gly337Glu</td>
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**Abbreviations:** LOD, linkage disequilibrium; SNP, single nucleotide polymorphism.

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30% of the $\varepsilon 3/4$ carriers suffered from LOAD due to their APOE genotype. Thus, it is highly unlikely that APOE $\varepsilon 3/4$ accounts for the heavy burden of AD in this family.

**TTC3** is an appealing LOAD candidate gene because of its involvement in pathways implicated in LOAD; functional studies in neuronal cell lines have shown that TTC3 plays an important role in ubiquitination, Akt signaling, negative cell cycle control, and inhibition of neuronal differentiation.\(^{29-31}\) In addition, the TTC3 protein carries conserved domains predicted to be involved in microtubule and actin-related processes of the cytoskeleton, apoptosis, cellular stress response, and mitochondrial function.\(^{32-36}\) Therefore, **TTC3** might act in multiple LOAD-relevant pathways. Moreover, these processes are intertwined with each other with respect to normal neuronal and synaptic functioning and Aβ and protein tau pathology.\(^{32,37-39}\) The identification of additional AD families with rare, nonsynonymous variants potentially contributing to disease would strengthen **TTC3** as an AD candidate gene.

During human embryogenesis, **TTC3** becomes predominantly expressed in neuronal tissue with high expression in cortical layers of the brain.\(^{40,41}\) Human postmortem brain microarray data demonstrate that cortical **TTC3** expression is consistently lower in LOAD cases compared to controls without dementia.\(^{2,3}\) Moreover, **TTC3** expression is negatively correlated with LOAD neuropathology markers Braak stage and frontal atrophy in the prefrontal cortex of LOAD cases. This pattern suggests that **TTC3** activity might protect against LOAD; thus, variants that reduce **TTC3** expression could contribute to LOAD risk. The p.S1038C substitution may alter the TTC3 protein’s conformation and thereby its function via the formation of bisulfide bonds with other cysteine residues, for instance in the short Cys-His-rich motif of the C-terminal H2 ring finger domain of the protein.\(^{42-45}\) A domain required for ubiquitin ligase activity,\(^{30}\) **TTC3** is an E3 ubiquitin ligase that acts as a specific terminal regulator of Akt signaling, a core intracellular pathway for cell survival.\(^{30}\) Akt/mTOR (mammalian target of rapamycin) signaling is a major cellular senescence pathway implicated in several age-related human diseases. Moreover, a large body of evidence attributes Akt/mTOR signaling as a key neuronal pathway that is disregulated in LOAD.\(^{46}\) Therapeutic approaches designed to normalize the Akt/mTOR axis protected against AD pathology and cognitive decline in animal models.\(^{47,48}\) Some of these therapies modifying this pathway in LOAD patients have been successful.\(^{49}\) Therefore, genetic variants that damage TTC3 protein function could contribute to LOAD susceptibility by reducing **TTC3**’s ability to counteract activated Akt, thereby facilitating an increase in activity of the Akt/mTOR axis. Further molecular studies that decipher the potential consequences of the alteration reported here are required to fully understand how the genetic alteration may contribute to AD pathogenesis.

Therefore, this study demonstrates how WES of single, extended families with a high burden of LOAD can lead to the identification of rare, predicted damaging genetic variants with potentially considerable risk to LOAD.

**AUTHOR CONTRIBUTIONS**


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and receives royalty payments from Duke University for CMT2A with Athena. Dr. Cuccaro has served on the editorial board of Child Psychiatry & Human Development. Dr. Gilbert has received research support from the Alzheimer Association. Dr. Schellenberg has served on the Alzheimer's Association Veterans Affairs Cooperative Study Unit Executive Committee, the Autism Genome Project, the National Alliance for Autism Research Medical Advisory Board, the Society of Progressive Supranuclear Palsy Scientific Advisory Board, and the Alzheimer Research Consortium Executive Board—Autism Genome Project Pfeiffer PSP Research Foundation; has received travel/speaker honoraria from the Alzheimer's Disease Center Directors Meeting, the CurePSP International Research Symposium, the Robert Katzman Memorial Lecture, the Colorado 25th Annual Southern California Alzheimer’s Disease Research Conference, the University of California Institute for Memory Impairment and Neurological Disorders, Biomarkers in Neuropsychiatric Disorders, NIH, Novartis Institutes for Biomedical Research, Inc., McKnight Brain Institute, Keep Memory Alive Event Center, Lou Ruvo Center for Brain Health, Accelerated Medicines Program Alzheimer’s Disease F2F Target Discovery and Preclinical Validation Project Meeting, and the American Association of Neuropathologists; has served on the editorial boards of the Journal of Neural Transmission, Alzheimer's Research, American Journal of Alzheimer's Disease & Other Dementias, Neurodegenerative Disease, Current Alzheimer Research, and Pathology and Laboratory Medicine International; is an employee of the University of Pennsylvania; has been a consultant for Finnigan Law Firm; and has received research support from Integregen, NIA/NIH, and CurePSP CBD Solutions. Dr. Carney and Dr. Pericak-Vance report no disclosures. Go to Neurology.org/ng for full disclosure forms.

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