

RESEARCH ARTICLE

Whole genome-wide sequence analysis of long-lived families (Long-Life Family Study) identifies *MTUS2* gene associated with late-onset Alzheimer's disease

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Abstract

INTRODUCTION: Late-onset Alzheimer's disease (LOAD) has a strong genetic component. Participants in Long-Life Family Study (LLFS) exhibit delayed onset of dementia, offering a unique opportunity to investigate LOAD genetics.

METHODS: We conducted a whole genome sequence analysis of 3475 LLFS members. Genetic associations were examined in six independent studies ($N = 14,260$) with a wide range of LOAD risk. Association analysis in a sub-sample of the LLFS cohort ($N = 1739$) evaluated the association of LOAD variants with beta amyloid ($A\beta$) levels.

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1 | BACKGROUND

Alzheimer's disease (AD) affects an estimated 50 million people worldwide. Approximately 90% of cases are sporadic and occur after 65 years of age, that is, late-onset AD (LOAD).¹ LOAD is complex and multifactorial with genetic and environmental factors² contributing to the brain accumulation of beta amyloid (A β) and tau proteins.³ Twin studies have demonstrated the high heritability of LOAD, with overall estimates ranging from 58% to 79%.⁴ Genome-wide association studies (GWAS) revealed that LOAD risk is driven by multiple loci. These GWAS genes confer only weak to modest AD risk when compared with a 4- to 15-fold increase in AD risk due to apolipoprotein E (APOE)- ϵ 4 inheritance. The identified susceptibility loci explain a small proportion of the LOAD indicating that undiscovered loci remain.^{5,6}

The largest GWAS meta-analysis to date⁷ identified 75 variants using genetic data from 880,000 subjects. Instead of identifying LOAD risk factors using large samples of the general population, we focused our discovery efforts on the Long-Life Family Study (LLFS). LLFS aims to examine the genetic and non-genetic factors associated with exceptional longevity. The LLFS cohort appears to be healthier than random cohorts of the same age/sex.⁸ Previous studies examining the health of LLFS participants⁹ have shown that offspring of long-lived family members exhibit lower rates of diabetes mellitus, pulmonary disease,

RESULTS: We identified several single nucleotide polymorphisms (SNPs) in tight linkage disequilibrium within the *MTUS2* gene associated with LOAD (rs73154407, $p = 7.6 \times 10^{-9}$). Association of *MTUS2* variants with LOAD was observed in the five independent studies and was significantly stronger within high levels of A β 42/40 ratio compared to lower amyloid.

DISCUSSION: *MTUS2* encodes a microtubule associated protein implicated in the development and function of the nervous system, making it a plausible candidate to investigate LOAD biology.

KEYWORDS

genetic risk, late-onset Alzheimer's disease, microtubule protein, *MTUS2* gene, whole genome sequence

Highlights

- Long-Life Family Study (LLFS) families may harbor late onset Alzheimer's dementia (LOAD) variants.
- LLFS whole genome sequence analysis identified *MTUS2* gene variants associated with LOAD.
- The observed LLFS variants generalized to cohorts with wide range of LOAD risk.
- The association of *MTUS2* with LOAD was stronger within high levels of beta amyloid.
- Our results provide evidence for *MTUS2* gene as a novel LOAD candidate locus.

and peripheral arterial disease. Consistent with previously reports,¹⁰ rates of dementia and memory decline among offspring of parents with exceptional longevity¹⁰ are lower, and cognitive tests scores in the LLFS offspring generation are higher on average when compared to spouse controls.¹¹ LLFS results demonstrated that individuals who are part of exceptionally long-lived families are protected against cognitive impairment characteristic of LOAD.¹² The genetic advantage is likely due to variants that decrease risk for aging-related diseases. The exceptionally healthy LLFS population provides a unique opportunity to investigate the genetic contributions to LOAD. Our previous research has also shown that reduction of the frequency of APOE- ϵ 4 allele and the increase in the frequency of the APOE- ϵ 2 allele in the LLFS offspring generation contribute to longevity.¹³

The majority of the genetic studies in LOAD have consisted of GWAS based on genotyping single nucleotide polymorphism (SNP) arrays or a combination of microarray-based genotyping with high dense imputation efforts.^{6,14-23} However, decreasing costs and advances in analytic methods have made high-throughput sequencing a feasible alternative.²⁴ Whole genome sequence (WGS) GWAS (Seq-GWAS) offers a significant advantage with the potential to detect all variants present in a sample, not only those present on an array or imputation reference panel. However, to date, those few Seq-GWAS studies of LOAD have either exclusively focused on rare variants²⁵ or specific populations.²⁶

The present study aims to: (1) examine the association between WGS variants and LOAD in the LLFS cohort, and (2) validating the findings of LLFS in additional cohorts with different risk of dementia: high risk populations (i.e., familial LOAD and Down's syndrome [DS]), AD referral-base cohort, and population-based cohorts.

The selection of the DS datasets as validation cohorts is based on its strong association with AD.²⁷ This association has a genetic basis through a gene-dose effect of the triplication of amyloid precursor protein (*APP*) gene in this population. Ninety percent of individuals with DS will have developed AD by age 70 (vs. 11% of people over age 65 in the general population). Because of these similarities, DS has been conceptualized as genetically determined AD, just like the autosomal dominant forms.

In the case of the familial LOAD, the most significant LOAD risk factor after advanced age is family history. The observed incidence rates for LOAD are estimated to be three to five times higher in multiplex families.²⁸ Individuals with a first-degree relative affected by AD, are at 4- to 10-fold higher risk for developing LOAD.²⁹⁻³²

2 | METHODS

2.1 | Study cohorts

In addition to LLFS, the present study examined the findings in six external datasets. These cohorts include varying LOAD risk, ranging from those with high risk (The National Institute on Aging Alzheimer's Disease Family Based Study [NIA-LOAD FBS], The Alzheimer Biomarkers Consortium—Down Syndrome [ABC-DS], and The Multiomic Studies of Alzheimer's Disease in Adults with Down Syndrome Study [omicsADDS]), as well as those from the general population (the Religious Orders Study/Memory and Aging Project [ROSMAP], the Alzheimer's Disease Genetics Consortium [ADGC], and the Washington Heights/Inwood Columbia Aging Project [WHICAP]).

2.1.1 | The Long-Life Family Study (LLFS)

The LLFS is a longitudinal, family-based study designed to assess genetic and environmental risk factors associated with exceptional longevity. Study design details can be found elsewhere.⁸ AD status was determined by a dementia review committee. Briefly, LLFS participants were selected for dementia review based on the presence of any of the following: (i) Clinical Dementia Rating³³ score greater than zero, (ii) cognitive impairment consistent with dementia using a previously published diagnostic algorithm,¹² and (iii) if previous data were missing, the informant report of a cognitive problem was used. Participants included in the current analyses included those diagnosed with probable LOAD with and without stroke. Participants with mild cognitive impairment (MCI) and other types of dementia were not included in the analyses. Additionally, Danish participants were excluded due to the lack of clinical consensus dementia diagnosis.

RESEARCH IN CONTEXT

- 1. Systematic review:** Late-onset Alzheimer's disease (LOAD) susceptibility loci identified explain a small proportion of its heritability, indicating that undiscovered loci remain. We focused on the Long-Life Family Study (LLFS), a cohort designed to assess genetic and environmental factors associated with exceptional longevity. The exceptionally healthy LLFS population provides a unique opportunity to investigate LOAD genetics. Our study identified *MTUS2* gene's variants associated with LOAD. The *MTUS2* association with LOAD was observed in six independent studies with a wide range of LOAD risk and was significantly stronger within high plasma levels of beta amyloid.
- 2. Interpretation:** *MTUS2* gene encodes a microtubule associated protein implicated in nervous system development and function, making it a plausible candidate for LOAD.
- 3. Future directions:** Future studies may seek to conduct functional validation analyses. Additionally, future studies may aim to consider the contribution of socioeconomic status, mental/behavioral health, and environmental factors to LOAD risk.

Whole genome sequencing was carried out by the McDonnell Genome Institute (MGI) at Washington University by Illumina Sequencers. MGI subsequently quality control procedures included: (i) alignment to build GRCh38 with BWA-MEM; (ii) marking duplicates with Picard; (iii) base quality score recalibration with GATK; (iv) loss-less conversion to CRAM format with SAMtools. The quality of the derived CRAM files was reviewed using to filter out samples with high level of contamination and haploid coverage. Final analysis files excluded SNP variants by coverage, lack of informativeness (monomorphic or missing), and failing Hardy-Weinberg test because of excess heterozygosity.

Plasma was collected in a random sample of 1739 LLFS participants, demographically comparable to the entire sample, and their corresponding $A\beta$ levels were processed after 24–48 h. Centrifuged plasma aliquoted in polypropylene tubes and stored at -80°C was used to measure $A\beta_{42}$ and $A\beta_{40}$ using Luminex technology.

2.1.2 | The National Institute on Aging Alzheimer's Disease Family Based Study (NIA-LOAD FBS)

The NIA-LOAD FBS is the largest collection of multiplex LOAD families recruited and longitudinally assessed worldwide. A detailed description of the study can be found elsewhere.²⁸ The present study used data from 4079 study participants. For analysis purposes, LOAD cases were defined using the National Institute of Neurological and

Communicative Disorders and Stroke and the Alzheimer's Disease and related disorders association (NINCDS-ADRDA) criteria.³⁴ Nondemented subjects were defined as any individual with no evidence of LOAD, and subjects with MCI were excluded from analyses. Genome-wide microarray data³⁵ were imputed using the HRC panel through the Michigan Imputation Server.³⁶

2.1.3 | The Alzheimer Biomarkers Consortium-Down Syndrome (ABC-DS) and The Multiomic Studies of Alzheimer's Disease in Adults with Down Syndrome Study (omicsADDS)

The ABC-DS is a multisite longitudinal study examining biomarkers of AD in a large cohort of adults with DS.³⁷ The omicsADDS study aims to identify multiomic profiles for successful cognitive aging in adults with DS. The recruitment and clinical assessment have been previously detailed.^{38–40} For both cohorts, the determination of diagnostic status was made during case consensus conferences. As previously described,³⁸ diagnostic status considered for the present analyses were (1) cognitively stable, (2) MCI-DS, (3) AD-dementia. To maximize our statistical power, subjects with MCI were grouped with cognitively stable.

Genome-wide genotyping in both cohorts was performed using the Illumina Infinium Global Screening Array-24 v2.0 at the Center for Applied Genomics at Children's Hospital of Pennsylvania. Imputation was performed using the TOPMed imputation server.³⁶ For the present analysis, the ABC-DS sample consisted of 366 participants and the omicsADDS consisted of 244 participants.

2.1.4 | The Religious Orders Study/Memory and Aging Project (ROSMAP)

Study participants were drawn from two different population-based cohorts: (i) The Religious Orders Study (ROS) study, which includes older Catholic nuns, priests, and brothers from groups across the United States and (ii) The Rush Memory and Aging Project (MAP), which includes older individuals from the metropolitan Chicago area. Detailed AD dementia are based on criteria of the joint working group of the National Institute of Neurological and Communicative Disorders and Stroke and the Alzheimer's Disease and Related Disorders Association (NINCDS/ADRDA).³⁴ Participants with MCI were excluded for the present analysis. As previously described, a subset of the ROSMAP samples ($n = 1200$) underwent whole genome sequencing.⁴² To minimize population structure bias, analyses were restricted to those with White ancestry, a total of 912 samples.

2.1.5 | The Alzheimer's Disease Genetics Consortium (ADGC)

The samples genotyped by the ADGC are distributed by the National Alzheimer's Coordinating Center (NACC). The NACC was established

by the National Institute on Aging in 1999 to facilitate collaborative research by using data collected from the approximately 30 NIA-funded Alzheimer's Disease Centers across the United States. Clinical diagnosis of Alzheimer's dementia is based on NINCDS-ADRDA criteria.³⁴ Participants classified as mild cognitive impaired were excluded for this current analysis. The genotyping of NACC samples was performed on a variety of genotyping platforms as previously described.²² Genetic imputation was carried out using IMPUTE v2 software⁴³ using the 1,000 Genomes Project as a reference.⁴⁴

2.1.6 | The Washington Heights/Inwood Columbia Aging Project (WHICAP)

Participants were drawn from a community-based multiethnic cohort via random sampling of Medicare beneficiaries living in northern Manhattan, New York.⁴⁵ Based on information obtained at the baseline and follow-up visits, dementia diagnosis was made by consensus during diagnostic conferences. The dementia diagnosis was based on Diagnostic and Statistical Manual of Mental Disorders⁴⁶ as well as NINCDS-ADRDA criteria.³⁴ Mild cognitive impaired participants were not included in the present analysis. As previously described,⁴⁷ genotypic data underwent imputation using IMPUTE2³⁶ with the HRC r1.1.2016 reference panel.

2.2 | Statistical analyses

SNP-based association analyses in LLFS and in NIA-LOAD FBS were conducted using generalized logistic mixed models implemented in GMMAT software.⁴⁸ To account for relatedness among participants in both cohorts, standardized relatedness matrix files were computed using GEMMA software.⁴⁹ Diagnosis of LOAD was used as outcome variable and sex, age, education, site of recruitment, kinship matrix, and first three principal components were modeled as covariates. In the DS cohorts, association analysis with *MTUS2* variants were performed using PLINK software.⁵⁰ Because of their intellectual disability, educational levels are not available for DS individuals. However, based on previous findings,⁵¹ all analyses have been adjusted for the severity of their intellectual disability. All models were adjusted for sex, age, level of functioning, and principal components. In ROSMAP, WHICAP, and ADGC cohorts, logistic models implemented in EPACTS software⁵² were used to investigate the association of *MTUS2* variants with LOAD. All analyses included sex, age, education, and principal components as covariates. In the LLFS cohort, SNPs were retained based on their minimum allele frequency ($MAF \geq 1\%$), Hardy-Weinberg p -value $\geq 5.7 \times 10^{-7}$, and genotype missing proportion $\geq 5\%$.

In LLFS, gene-based test of association for *MTUS2* gene were conducted using EPACTS⁵² software. Sex, age, education, and principal components adjusted analyses were conducted using the sequence kernel association test SKAT and incorporating the efficient Mixed-Model Association eXpedited (EMMAX) algorithm.^{49,53}

Plasma $A\beta$ levels ($A\beta_{40}$ and $A\beta_{42}$) were collected at visit two in a sub-sample of 1739 LLFS participants. The socio-demographic

TABLE 1 Characteristics of study cohorts.

	LLFS			Replication datasets					
	All	Proband	Offspring	NIA-FBS	ABC-DS	omicsADDS	ROSMAP	ADGC	WHICAP
N	3475	1303	2172	4079	336	244	912	6977	851
Percentage of women	55	53	56	61	46	65	66	57	60
Age (mean ± SD)	71 ± 16	89 ± 7	61 ± 8	74 ± 11	46 ± 10	56 ± 7	89 ± 6	79 ± 8	81 ± 7
Education (mean ± SD)	14 ± 3	13 ± 2	16 ± 3	14 ± 3	-	-	16 ± 4	16 ± 3	14 ± 4
Prevalence of LOAD (%)	7	15	2	60	21	28	69	48	21

Abbreviations: ABC-DS, Alzheimer's Biomarkers Consortium-Down Syndrome; ADGC, Alzheimer's Disease Genetic Consortium; LLFS, Long-Life Family Study; LOAD, late-onset Alzheimer's disease; NIA-FBS, The National Institute on Aging Alzheimer's Disease Family Based Study; omicsADDS, The Multi-omic Studies of Alzheimer's Disease in Adults with Down Syndrome Study; ROSMAP, the Religious Orders Study/Memory and Aging Project; WHICAP, the Washington Heights/Inwood Columbia Aging Project.

^aAs described in the manuscript, educational levels are not available for both Down Syndrome cohorts.

characteristics of the sample were comparable to the whole LLFS cohort (data not shown). Based on previous findings that have consistently reported that a lower $A\beta_{42/40}$ ratio in plasma is associated with higher risk of AD,⁵⁴⁻⁵⁷ we investigated whether the association of *MTUS2* with LOAD was influenced by $A\beta_{42/40}$ levels. Because of the relatively limited sample size, we speculated that association effects will be more evident using quartiles of $A\beta_{42/40}$ ratio. The $A\beta_{42/40}$ ratio quartiles were categorized as a dichotomous variable: quartiles 1 and 2 were considered as lowest quartiles; quartiles 3 and 4 were considered the highest quartiles. Association analyses were performed using generalized estimating equations in SPSS software to adjust for relatedness among LLFS's participants. We investigated the association of $A\beta_{42/40}$ ratio levels with the *MTUS2* variants identified in the GWAS analysis (11 SNPs in tight linkage disequilibrium [LD]), adjusting for sex, age, and LOAD diagnosis.

3 | RESULTS

Characteristics of the study's participants are summarized in Table 1. The proportion of women ranged from 46% in ABC-DS to 66% in ROSMAP. The average age of participants from the proband LLFS generation and ROSMAP was significantly older (89 ± 7 years and 89 ± 6 , respectively) when compared to the participants from the other cohorts. The highest prevalence of LOAD (69%) was observed for ROSMAP participants, followed by the NIA-LOAD FBS (60%). The highest educational level was observed for ROSMAP and ADGC with an average of 16 years of education.

In LLFS, association results from the SNP-based analysis are shown in Figures 1 and 2. Several SNPs in chromosome 1 reached genome-wide significance levels ($p \leq 5 \times 10^{-8}$); however, they cover a large genomic area of more than 500Kb, which encompassed at least four different genes, none of which reached significance at the gene level. Tables S1-S5 list the single-SNP association results for those chromosomal regions in which at least one SNP reached the genome-wide significance threshold (chromosomes 1, 6, 13, 20, and 21). In addition, Figures S1-S5 show the corresponding chromosomal regional associa-

tion plots using Locuszoom.^{58,59} On chromosome 13, a tight LD block of 11 SNPs spanning 1.6Kb (Table 2) was mapped to Microtubule Associated Scaffold Protein 2 gene, *MTUS2*. The strongest association was observed at *MTUS2* intronic variant rs73154407 ($p = 7.6 \times 10^{-9}$). Gene-based analyses also reached statistical significance, nominating *MTUS2* gene as a LOAD risk locus ($p = 2.6 \times 10^{-8}$). In secondary analyses considering adjustment for *APOE-ε4* allele, *MTUS2* associations remained statistically significant (data not shown). Likewise, we did not observe either a significant association of *APOE* locus with AD, consistent with our previous reports¹³ on the lower likelihood of carrying an *APOE-ε4* allele as well as an elevated allele frequency of *APOE-ε2*⁶⁰ among LLFS family members compared to similar-aged spouses' controls.

The association of *MTUS2* with LOAD observed in LLFS was validated using six different independent cohorts (Table 3). Due to the different genetic coverage of WGS and GWAS data, *MTUS2* associated SNPs differ across the cohorts, and therefore meta-analyses cannot be performed. We performed LD calculations using the NIH Web-based application LDlink (LD matrix module; <https://ldlink.nci.nih.gov/> = home). When assessing the LD between the top SNP in LLFS and SNPs in the replication cohorts, our results showed high LD measured as D' , but moderate-low values as measured by r^2 ($r^2 < 0.5$).

The strongest association in the NIA-LOAD FBS was observed at SNP variant rs775323299 ($p = 8.4 \times 10^{-5}$). Next, we analyzed two independent samples of DS. SNP variants rs4769747 and rs55646198 yielded the most significant association signal (ABC-DS $p = 0.009$ and omicsADDS $p = 0.009$, respectively). In ROSMAP, WHICAP, and ADGC, the strongest associations were achieved at SNP rs9551636 ($p = 4 \times 10^{-4}$), rs68054178 ($p = 3.7 \times 10^{-5}$), and rs581625 ($p = 0.002$), respectively.

Results from LLFS (Table S6, Figure S6) revealed that *MTUS2* variant rs73154415 was strongly associated with plasma $A\beta_{42/40}$ ratio ($B = 1.06$, $SE = 0.01$, $p = 1.9 \times 10^{-5}$). Secondary analyses (Figure S1) demonstrated that the association of variant rs73154415 with LOAD become significantly stronger within high $A\beta_{42/40}$ ratio compared to lower amyloid ratio ($B = 3.46$, $SE = 0.74$, $p = 3 \times 10^{-6}$ vs. $B = 1.66$, $SE = 0.67$, $p = 0.005$, respectively).

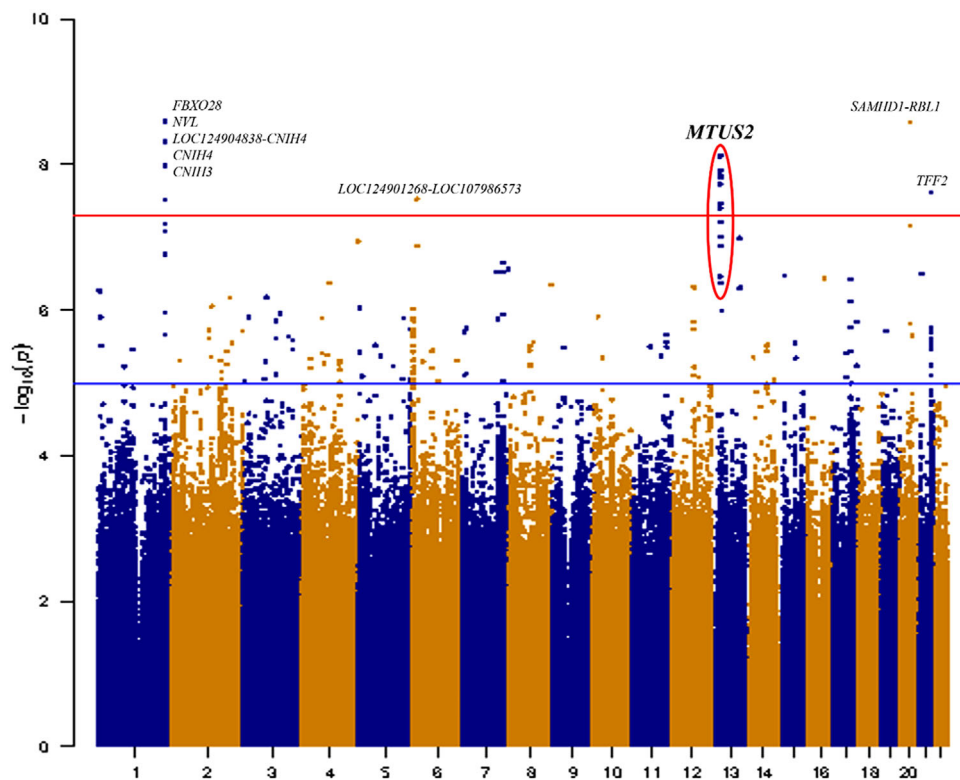


FIGURE 1 Manhattan plot of Seq-GWAS results in LLFS cohort. The X-axis represents the genomic position for each of the SNPs analyzed; Y-axis represents the $-\log_{10}$ transformed p -values. The red line indicates the genome-wide significance threshold ($p < 5 \times 10^{-8}$). GWAS, genome-wide association studies; LLFS, Long-Life Family Study; SNP, single nucleotide polymorphism.

TABLE 2 GWAS top significant *MTUS2* variants associated with late-onset Alzheimer's disease.

Chr	SNP	hg38	A1	A2	N	MAF	B	SE	p -Value
13	rs7330699	29,483,430	G	A	3,200	0.010	2.89	0.51	1.4×10^{-8}
13	rs73154407	29,483,753	G	T	3,184	0.010	2.97	0.51	7.6×10^{-9}
13	rs11842476	29,483,758	G	A	3,184	0.013	2.47	0.46	6.1×10^{-8}
13	rs73154410	29,483,772	A	T	3,187	0.010	2.96	0.51	7.8×10^{-9}
13	rs1350113738	29,483,860	T	C	3,184	0.010	2.88	0.51	1.5×10^{-8}
13	rs17073514	29,484,261	G	T	3,202	0.013	2.41	0.45	9.8×10^{-8}
13	rs116813765	29,484,326	C	T	3,204	0.010	2.89	0.51	1.4×10^{-8}
13	rs73154413	29,484,390	C	T	3,207	0.011	2.85	0.51	1.8×10^{-8}
13	rs73154414	29,484,568	A	G	3,201	0.010	2.91	0.51	1.2×10^{-8}
13	rs73154415	29,484,701	A	G	3,206	0.010	2.77	0.50	3.5×10^{-8}
13	rs41291229	29,484,987	A	G	3,186	0.010	2.75	0.50	4.0×10^{-8}

Note: Physical positions in base pair (bp) correspond to the hg38 genome build.

Abbreviations: GWAS, genome-wide association studies; MAF, minimum allele frequency; SNP, single nucleotide polymorphism.

4 | DISCUSSION

Genome-wide analysis of WGS data from healthy aging families identified *MTUS2* as a novel LOAD genetic risk factor. The association of *MTUS2* variants with LOAD was also observed in the six independent cohorts, including populations at high risk of LOAD. The association

became stronger among subjects with high levels of plasma $A\beta_{42/40}$ ratio.

MTUS2 is a plus end tracking protein involved in elongation of microtubules,⁶¹ expressed in brain during development and adulthood.⁶² Few studies reported variants in *MTUS2* associated with LOAD. In a genome-wide interaction study of cerebrospinal

TABLE 3 Results of *MTUS2* association with LOAD in independent cohorts.

Cohort	SNP	bp (hg37)	N	B	SE	<i>p</i>	MAF	D'
NIA-LOAD	rs775323299	28,795,404	4079	3.92	0.50	8.4×10^{-5}	1.5×10^{-4}	NA
ABC-DS	rs73164590	29,064,021	366	2.01	0.35	0.002	0.01	1.00
omicsADDS	rs55646198	29,486,171	244	1.08	0.24	0.009	0.06	1.00
ROSMAP	rs118034475	29,426,891	625	1.15	0.40	0.004	0.01	1.00
WHICAP_NHW	rs68054178	29,128,208	839	4.12	0.62	3.7×10^{-5}	0.07	1.00
ADGC	rs581625	28,956,466	6977	3.15	0.60	0.002	0.17	0.42

Abbreviations: ABC-DS, Alzheimer's Biomarkers Consortium-Down Syndrome; ADGC, Alzheimer's Disease Genetic Consortium; LOAD, late-onset Alzheimer's disease; NIA-LOAD, The National Institute on Aging Alzheimer's Disease-Late-Onset Alzheimer's disease; omicsADDS, The Multiomic Studies of Alzheimer's Disease in Adults with Down Syndrome Study; ROSMAP, the Religious Orders Study/Memory and Aging Project; SNP, single nucleotide polymorphism; WHICAP, the Washington Heights/Inwood Columbia Aging Project.

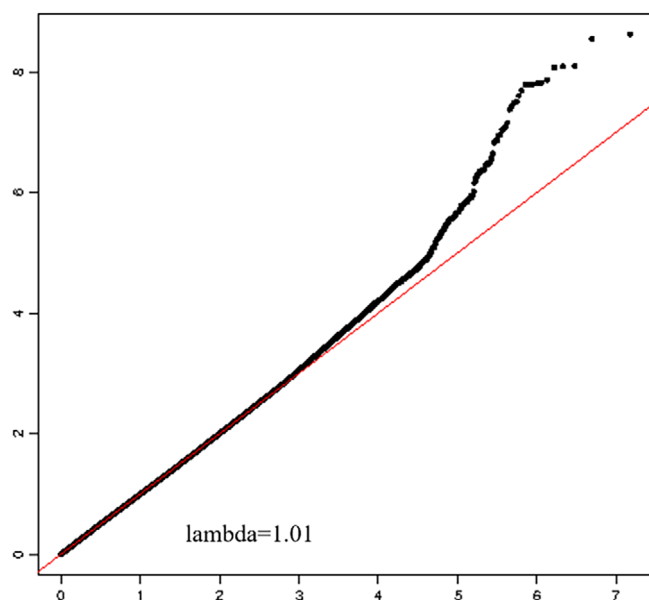


FIGURE 2 Q-Q plot of Seq-GWAS results in LLFS cohort. The X-axis represents the expected $-\log_{10}$ transformed *p*-values for each SNP association; the Y-axis represents the observed $-\log_{10}$ transformed *p*-values. GWAS, genome-wide association studies; LLFS, Long-Life Family Study; SNP, single nucleotide polymorphism.

fluid T-tau/ $A\beta_{42}$ ratio using the Alzheimer's Disease Neuroimaging Initiative cohort, *MTUS2* variants significantly interacted with *RLBP1L1* to explain a relatively high-level variance of this biomarker, demonstrating their potential association with LOAD pathology.⁶³ Variants in *MTUS2* were reported associated with an inflammatory LOAD subtype after having integrated *post mortem* brain gene co-expression data from LOAD samples.⁶⁴ Chen and colleagues,⁶⁵ nominated *MTUS2* as likely associated with AD using an epistasis detection method that completes a spatial search through a genetic algorithm. Results from the Accelerating Medicines Partnership for Alzheimer's disease (AMP-AD) consortium RNA-seq analysis of *post mortem* brains of more than 1100 individuals (<https://agora.adknowledgeportal.org/>), showed that *MTUS2* was significantly expressed across different brain regions (Figure S7). Moreover, three of the brain regions (cerebellum, parahippocampal gyrus, and temporal cortex) showed significant

change in *MTUS2* gene expression when comparing LOAD cases and cognitively healthy samples (Figure S8).

Our results suggest that *MTUS2* may play a role in $A\beta$ metabolism. $A\beta$ and tau protein deposition are the two principal pathological hallmarks of AD. The imbalance between $A\beta$ production and clearance leads to the aggregation of amyloid plaques, while hyperphosphorylated tau protein transform into neurofibrillary tangles.^{66,67} However, the relationship between amyloid and tau in the pathophysiology of LOAD remains unclear. It is known that tangles deposition is associated with cognitive decline,^{68,69} while mutations in the *APP* cause familial AD.⁷⁰ The current literature suggests that plaques and tangles may act synergistically as both are necessary for cognitive decline and glucose hypometabolism. Evidence in cellular and mouse models also shows a synergistic effect, with the presence of amyloid peptides increasing the formation of certain species of hyperphosphorylated tau as well as of its aggregates.^{71,72} The role of tau in amyloid plaque formation is less clear, although in at least one experiment, the presence of mutant tau increased the number of plaques in an *APP/PSEN1* mouse model.⁷³ In another study using tau-null animals, synapses were protected from $A\beta$ toxicity, highlighting the potential synergy between the two proteins.⁷⁴

The integrity of microtubules is essential for neurons to maintain their morphology and to transport components between cell body and synaptic terminals. $A\beta$ peptides can disrupt this transport through cytoskeleton reorganization which increases *APP* retention and its vesicular cleavage.⁷⁵ The mechanism through which $A\beta$ affects microtubule stability involves deetyrosination that causes hyperstabilization of the microtubules, which is a process dependent on *APP* expression. Although no direct link has been observed between *MTUS2* and tau phosphorylation or microtubule hyperstabilization, the fact that it is involved in microtubule dynamics could explain the association of genetic variations of this protein and LOAD. Indeed, the fact that our variant is associated with plasma $A\beta$ ratio supports the idea that microtubule stability is involved in *APP* metabolism.

Our study presents some limitations. First, differences in methodological approaches across the studies may introduce heterogeneity through biases variably, consequently affecting the results. Nonetheless, we have consistently shown an association between *MTUS2* gene and AD in all the studies, suggesting that our findings are robust and reliable. Second, we did not consider the contribution of additional

socioeconomic status, mental or behavioral health, and clinical comorbid conditions that may be associated with LOAD risk. Third, additional genetic factors in conjunction with lifestyle-behavior-environmental factors and their interactions may be needed for developing the disease. Fourth, functional analyses are needed to validate *MTUS2* as candidate risk gene.

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CONFLICT OF INTEREST STATEMENT

Bradley Christian receives PET precursor and compounds from Avid Radiopharmaceuticals Inc and equipment from Cerveau Technologies. Handen receives funding from National Institute of Child Health and Human Development, Autism Speaks, and Roche Pharmaceuticals. Mapstone is an inventor on patents related to biomarkers of neurodegenerative diseases owned by Georgetown University and the University of Rochester. The other authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest. Author disclosures are available in the [supporting information](#).

CONSENT STATEMENT

All study participants provided written informed consent and the study procedures were approved by the institutional review boards within

each institution. All study procedures were performed in accordance with the Declaration of Helsinki ethical principles.

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SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

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