## RESEARCH

**Open Access** 

# Large multi-ethnic genetic analyses of amyloid imaging identify new genes for Alzheimer disease



Muhammad Ali<sup>1,2†</sup>, Derek B. Archer<sup>3†</sup>, Priyanka Gorijala<sup>1,2</sup>, Daniel Western<sup>1,2</sup>, Jigyasha Timsina<sup>1,2</sup>, Maria V. Fernández<sup>1,2</sup>, Ting-Chen Wang<sup>3</sup>, Claudia L. Satizabal<sup>4,6,12</sup>, Qiong Yang<sup>5</sup>, Alexa S. Beiser<sup>5,6,12</sup>, Ruiqi Wang<sup>7</sup>, Gengsheng Chen<sup>8,9</sup>, Brian Gordon<sup>8,9</sup>, Tammie L. S. Benzinger<sup>8,9</sup>, Chengjie Xiong<sup>8</sup>, John C. Morris<sup>8,10</sup>, Randall J. Bateman<sup>8,10,22</sup>, Celeste M. Karch<sup>1</sup>, Eric McDade<sup>10</sup>, Alison Goate<sup>11</sup>, Sudha Seshadri<sup>12,13</sup>, Richard P. Mayeux<sup>14</sup>, Reisa A. Sperling<sup>15,16</sup>, Rachel F. Buckley<sup>16,17</sup>, Keith A. Johnson<sup>18</sup>, Hong-Hee Won<sup>19</sup>, Sang-Hyuk Jung<sup>19</sup>, Hang-Rai Kim<sup>20</sup>, Sang Won Seo<sup>21</sup>, Hee Jin Kim<sup>19,21</sup>, Elizabeth Mormino<sup>22</sup>, Simon M. Laws<sup>23</sup>, Kang-Hsien Fan<sup>24</sup>, M. Ilyas Kamboh<sup>24</sup>, Prashanthi Vemuri<sup>25</sup>, Vijay K. Ramanan<sup>26</sup>, Hyun-Sik Yang<sup>27,28,29,30</sup>, Allen Wenzel<sup>31</sup>, Hema Sekhar Reddy Rajula<sup>32</sup>, Aniket Mishra<sup>32</sup>, Carole Dufouil<sup>32</sup>, Stephanie Debette<sup>32,33,34</sup>, Oscar L. Lopez<sup>35</sup>, Steven T. DeKosky<sup>36</sup>, Feifei Tao<sup>37</sup>, Michael W. Nagle<sup>37</sup>, Knight Alzheimer Disease Research Center (Knight ADRC), the Dominantly Inherited Alzheimer Network (DIAN), Alzheimer's Disease Neuroimaging Initiative (ADNI), ADNI-DOD, A4 Study Team, the Australian Imaging Biomarkers, Lifestyle (AIBL) Study, Timothy J. Hohman<sup>3</sup>, Yun Ju Sung<sup>1,2</sup>, Logan Dumitrescu<sup>3†</sup> and Carlos Cruchaga<sup>1,2,8,38,39\*†</sup>

## Abstract

Amyloid PET imaging has been crucial for detecting the accumulation of amyloid beta (Aβ) deposits in the brain and to study Alzheimer's disease (AD). We performed a genome-wide association study on the largest collection of amyloid imaging data (N = 13,409) to date, across multiple ethnicities from multicenter cohorts to identify variants associated with brain amyloidosis and AD risk. We found a strong *APOE* signal on chr19q.13.32 (top SNP: *APOE*  $\epsilon$ 4; rs429358;  $\beta$ =0.35, SE=0.01, P=6.2 × 10<sup>-311</sup>, MAF=0.19), driven by *APOE*  $\epsilon$ 4, and five additional novel associations (*APOE*  $\epsilon$ 2/rs7412; rs73052335/rs5117, rs1081105, rs438811, and rs4420638) independent of *APOE*  $\epsilon$ 4. *APOE*  $\epsilon$ 4 and  $\epsilon$ 2 showed race specific effect with stronger association in Non-Hispanic Whites, with the lowest association in Asians. Besides the *APOE*, we also identified three other genome-wide loci: *ABCA7* (rs12151021/chr19p.13.3;  $\beta$ =0.07, SE=0.01, *P*=9.2 × 10<sup>-09</sup>, MAF=0.32), *CR1* (rs6656401/chr1q.32.2;  $\beta$ =0.1, SE=0.02, P=2.4 × 10<sup>-10</sup>, MAF=0.18) and *FERMT2* locus (rs117834516/chr14q.22.1;  $\beta$ =0.16, SE=0.03, *P*=1.1 × 10<sup>-09</sup>, MAF=0.06) that all colocalized with AD risk. Sex-stratified analyses identified two novel female-specific signals on chr5p.14.1 (rs529007143,  $\beta$ =0.79, SE=0.14, P=1.4 × 10<sup>-08</sup>, MAF=0.006, sex-interaction P=9.8 × 10<sup>-07</sup>) and chr11p.15.2 (rs192346166,  $\beta$ =0.94, SE=0.17, P=3.7 × 10<sup>-08</sup>, MAF=0.004, sex-interaction P=1.3 × 10<sup>-03</sup>). We also demonstrated that the overall genetic architecture

<sup>†</sup>Muhammad Ali, Derek B Archer are equal co-first authors and Logan Dumitrescu and Carlos Cruchaga equal co-last authors for this article

\*Correspondence: Carlos Cruchaga cruchagac@wustl.edu Full list of author information is available at the end of the article



© The Author(s) 2023. **Open Access** This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/ficenses/by/4.0/. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated in a credit line to the data.

of brain amyloidosis overlaps with that of AD, Frontotemporal Dementia, stroke, and brain structure-related complex human traits. Overall, our results have important implications when estimating the individual risk to a population level, as race and sex will needed to be taken into account. This may affect participant selection for future clinical trials and therapies.

Keywords Brain amyloidosis, Amyloid PET, Alzheimer's disease, Multi-ethnic, Meta-analysis, GWAS

## Introduction

Alzheimer's disease (AD) is a complex polygenic disease with a genetic heritability estimated to be 58–79% [1]. This high genetic heritability in AD provides the opportunity to perform large-scale genetic studies in order to characterize new biological features, identify relevant pathophysiological processes, and establish novel diagnostic biomarkers for early detection. Recent genomewide association studies (GWAS) have identified more than 74 AD risk loci, including the *APOE* e4 locus, implicating various biological process e.g., amyloid processing and innate immunity in the development of AD [2–5]. However, known common AD variants account only for an approximately 30% of the AD genetic variance [6] and a large proportion of the underlying heritability still remains unexplained.

Although most of the genetic studies on AD are focused on clinical diagnosis as the primary outcome, using quantitative endophenotypes can also be helpful in identifying additional AD-related genes. Two such ADrelated endophenotypes are accumulation of amyloidbeta (A $\beta$ ) in the brain and the formation of tau deposits in the form of neurofibrillary tangles and dystrophic neurites (tau pathology) [7]. Different cross-sectional and longitudinal studies on cognitively normal subjects have also implicated amyloidosis as an early process in AD pathology [8–10]. The *in-vivo* detection of A $\beta$  accumulation in the brain, as measured by positron emission tomographic (PET) imaging tracers such as 11C-labeled Pittsburgh Compound-B (PiB) [11] and 18F-AV-45 (Florbetapir) [12] has provided a biomarker for AD diagnosis and risk assessment. Development of this advanced brain imaging approach has enabled the detection of fibrillar A $\beta$  before the onset of symptoms, providing avenues for characterizing new genetic risk factors, and to design novel therapeutic approaches to halt the early progression of disease.

Previous genetic investigations leveraging amyloid PET imaging data as AD endophenotype have established its association with *APOE* locus [11–14]. In a recent multicenter case–control based study (N=4314) using amyloid PET as a quantitative trait [15], a novel locus was reported to be associated with brain amyloidosis within *RBFOX1* gene. However, clinical heterogeneity across these studies and their limited sample size demands

further investigations to replicate and expand on these findings. Here, we systematically analyzed the largest collection of amyloid imaging data (N = 13,409), across multiple ethnicities from multicenter cohorts (Knight ADRC, A4, DIAN, ADNI, ADNIDOD, UPitt, HABS, AIBL, Memento, MCSA, WRAP, Berkeley, Korean study, and MISSION-AD) as a quantitative trait to identify the functional variants and genes driving the association of AD. Furthermore, we have conducted gender-, and *APOE*stratified analyses to investigate the effect of these variables on brain amyloidosis.

## Methods

## Study samples and amyloid-PET harmonization

For this study, we collected data from 14 different cohorts with a total sample size of 13,409 participants (Table 1). We analyzed the association of common and low frequency (MAF>0.0005) genetic variants with amyloid-PET imaging, which is a well-known and validated AD endophenotype, serving as biomarker for brain amyloidosis. These subjects were recruited from the Memory and Aging Project (MAP) at the Knight Alzheimer Disease Research Center (Knight-ADRC) [16, 17], Alzheimer's Disease Neuroimaging Initiative (ADNI) [18], the Dominantly Inherited Alzheimer Network (DIAN) [19], Anti-Amyloid Treatment in Asymptomatic Alzheimer's Disease (A4) [20], ADNI Department of Defense (ADNIDOD) studies, Australian Imaging, Biomarkers and Lifestyle (AIBL) [21], The Harvard Aging Brain Study (HABS) [22], and University of Pittsburgh (UPitt) [23]. Furthermore, summary statistics data was obtained from six additional cohorts that processed the raw genotype and phenotype data according to the same pipeline. These cohorts included Memento [24], Mayo Clinic Study of Aging (MCSA) [25], Wisconsin Registry for Alzheimer's Prevention (WRAP) [26], Berkeley Aging Cohort study (BACS) [27], Korean study [28], and MIS-SION-AD. Part of the data used in the preparation of this article was obtained from the ADNI database (adni. loni.usc.edu). Collection of genotype data and PET image processing for each cohort are described in detail in the respective studies [16-23, 25, 27-29].

Briefly, individuals were diagnosed as cognitively healthy (controls) or clinical AD (cases), based on their Clinical Dementia Rating<sup>®</sup> ( $CDR^{®}$ ) that was available for

normalized Disease Rest Alzheimer's Pittsburgh; N Florbetapir; I	individua earch Cer Disease; A MCSA, May PIB, Pittsbi	ally to acc nter; ADN ADNI-DOE iyo Clinic : urgh Com	count for II, Alzheir D, ADNI D Study of , npound-l	normalized individually to account for within cohort Disease Research Center; ADNI, Alzheimer's Disease Ne Alzheimer's Disease; ADNI-DOD, ADNI Department of D Pittsburgh; MCSA, Mayo Clinic Study of Aging; WRAP, Wi Florbetapir; PIB, Pittsburgh Compound-B; FMT, Flutemet	variation. Abb euroimaging Ir befense studies 'isconsin Regist tamol	oreviatio nitiative; s; AIBL, A try for Al:	ns: PET, <sub>F</sub> DIAN, th <sub>i</sub> ustralian l zheimer's	normalized individually to account for within cohort variation. Abbreviations: PET, positron emission tomography; sd, standard deviation; Knight-ADRC, Knight Alzheimer's Disease Research Center; ADNI, Alzheimer's Disease Neuroimaging Initiative; DIAN, the Dominantly Inherited Alzheimer Network; A4, Anti-Amyloid Treatment in Asymptomatic Alzheimer's Disease; ADNI-DOD, ADNI Department of Defense studies; AIBL, Australian Imaging, Biomarkers and Lifestyle; HABS, The Harvard Aging Brain Study; UPitt, University of Pittsburgh; MCSA, Mayo Clinic Study of Aging; WRAP, Wisconsin Registry for Alzheimer's Pack, Berkeley Aging Cohort study; SUVR, standardize uptake value ratios; FBP, Fiorbetapir; PIB, Pittsburgh Compound-B, FMT, Flutemetamol	in tomo <u>c</u> herited A kers and .S, Berkelu	Jraphy; sd, Alzheimer N Lifestyle; H ey Aging C	standard de Network; A4, IABS, The Har Cohort study;	eviation; K Anti-Amy vard Agin SUVR, star SUVR, star	้(night-AD loid Treat g Brain St าdardize เ	RC, Knigl ment in <i>i</i> udy; UPit uptake va	it Alzheimer's symptomatic , University of ue ratios; FBP,
Variables	Total	A4	ADNI	AIBL	ADNI-DOD	DIAN	HABS	Knight-ADRC	UPitt	Korean	Memento	MCSA	WRAP	BACS	MISSION-AD
Tracer	I	SUVR	FBP	FBP/PIB/FMT	AV45	PIB	PIB	PIB/AV45	PIB	AV45	FBP/FMT	PIB	PIB	PIB	FBP/FBB/ FMT
Total	13,409	3,180	1,134	1,214	169	209	258	1,048	345	966	630	1712	78	173	2,263
Female (%)	48.52	60.09	47	54.04	0.59	52.63	59.3	54.77	45.8	56.9	59.25	46.3	35	59	48.6
Male (%)	51.48	39.91	53	45.96	99.41	47.37	40.7	45.23	54.2	43.1	40.75	53.7	65	41	51.4
Age (mean)	69.63	71.30	73.73	72.65	69.08	36.60	73.99	70.74	78.39	70.15	71.55	73.4	67	74.49	71.8
Age (sd)	7.65	4.73	7.62	6.62	4.67	10.86	6.1	9.01	9.91	8.72	8.15	10.6	6.02	6.30	7.8
APOE4 + (%)	35.06	36.04	43.39	35.01	26.63	28.71	29.84	41.13	32.46	41.7	28.65	28.6	42	28	48.7
Cases (%)	35.17	0.06	61.55	7.5	32.54	45.01	27.13	16.98	12.17	69.8	100	15.7	5	0	66
Controls (%)	60.69	99.94	32.36	79.74	67.46	38.28	72.87	61.83	87.83	30.2	0	84.1	95	100	0

e4-carriers (*APOE* e4 + participants), and percentage of cases and control participants, where available. To normalize amyloid PET endophenotype across different cohorts, we converted different amyloid imaging measures (e.g., Centiloid, PIB, and AV45) into log-normalized z-score using "scale" function in base R. Phenotype from each cohort was For each cohort, we report number of participants, percentage of females and males, mean age of the participants and standard deviation (SD) in the age, percentage of APOE Table 1 Demographics of amyloid PET GWAS participants at the time of scanning. This table summarizes basic demographic information of participants included in the analysis.

86% of the complete dataset. The CDR is a scaling system that categorizes the overall dementia severity for each participant into five classes (no dementia=0, very mild=0.5, mild=1, moderate=2, and severe=3). For this study, individuals with CDR=0 were considered as controls and the remaining were classified as cases. Participants were included if they had measurements of raw amyloid-PET levels and corresponding genotype data. Any participant that was missing information about the sex, age, and genetic principal components (PCs), was excluded from the study.

For standardization, amyloid PET data from each cohort was normalized to their reference cerebellar regions in order to obtain standardized uptake value ratios (SUVR) in a composite of cortical brain areas. As different cohorts obtained quantitative amyloid PET data using different tracers (e.g., PIB, FBP, and AV45), the available raw phenotypic data cannot combine unless it is normalized in advance. To normalize amyloid PET endophenotype across different cohorts, we converted different amyloid imaging measures into log-normalized z-score using "scale" function in base R. Briefly, z-scores were calculated by using the mean and standard deviation (SD) units across each cohort and applied to the entire endophenotype in order to account for within cohort variation. Samples having normalized z-score 3-SD away from the mean of the population were considered as outliers and removed from the subsequent analyses.

For seven additional cohorts (Memento, MCSA, WRAP, BACS, Korean study, and MISSION-AD), the raw phenotype and genetic data was not accessible due to the strict patient data sharing policies (Table 1). In those particular cases, the summary statistics data was obtained where association analyses were performed using the same analytical pipeline.

## Genotyping, imputation, quality control, and population structure

The genotyping platforms used by each cohort are listed in the respective studies [16–23, 25, 27–29]. All the GWAS datasets were aligned to GRCh38. For phasing and imputation of non-genotyped single-nucleotide polymorphisms (SNPs), we used the TOPMed Imputation Server (https://imputation.biodatacatalyst.nhlbi.nih.gov/#!). Phasing was performed using eagle v2.4 [30] and only those variants having imputation quality (Rsq or estimated R<sup>2</sup>) of 0.3 or greater were retained [31]. Genotyped and imputed variants with minor allele frequency (MAF) < 0.0002 were removed. We applied stringent quality control (QC) filters to process the genotyping array and sequencing data. We used the threshold of 98% for removing single nucleotide polymorphisms (SNPs) and individuals with low call rate. Autosomal

SNPs that were not in the Hardy–Weinberg equilibrium  $(P=1\times10^{-6})$  were also removed. Subject duplication and relatedness were estimated from identity-by-descent (IBD) analysis carried out in Plink version 2.0 [32]. In case of related subjects (Pihat $\geq$ 0.25), the sample from the Knight ADRC or with a higher number of variants that passed the QC was prioritized. We performed Principal component analysis (PCA) on the genotype data to obtain genetic PCs that capture population substructure. In order to evaluate the association of genetic variants with brain amyloidosis in different populations, we considered a homogenous pool of three ethnicities (Non-Hispanic whites, American African, and Asian) for the subsequent statistical analyses (Additional file 1: Fig. S1).

After QC of genotype and phenotype data, a total of 7557 participants remained available from eight different cohorts for the subsequent analyses. According to the genetic PCA, participants were determined to be Non-Hispanic Whites (NHW; n = 7036), American Africans (AFR; n = 359), and Asians (ASN; n = 162). Additionally, summary statistics data was leveraged from six external amyloid PET cohorts to further increase the study sample size for NHW (n = 11,556) and ASN (n = 1494) ancestries (Additional file 1: Table S1a and S1b). The demographic characteristics of all the study participants (n = 13,409)from each of the 14 datasets included in the multi-ethnic standard error (StdErr)-based meta-analysis are shown in Table 1. The distribution of standardized amyloid-PET levels (Z-score) across eight cohorts is shown in Additional file 1: Fig. S2. Overall, the participants ranged from age 37 to 78 with a mean age of 69 across all cohorts. The age of participants from the DIAN cohort was almost half the mean age of all cohorts and nearly all the samples in the ADNI-DOD cohort were males. Around onethird (34%) of the individuals were APOE  $\varepsilon$ 4 carriers. The number of cases and controls samples were 23% and 63%, respectively, and 14% of the individuals were missing the clinical diagnosis. There was an even distribution of gender across all cohorts with a total of 48% female and 52% male participants.

## Statistical analyses

Statistical analyses and data visualization were performed in Plink version 2.0 [32], Metal [33], and R version 3.5.2 [34]. We performed association analyses of common and low frequency variants with quantitative AD endophenotype from PET scan (A $\beta$ ) using an additive model and included sex, age and the 10 principal components as covariates, within each ethnicity. Within each ethnicity we first performed joint analyses using the Z-scores for those cohorts for which we had access to the genetic and amyloid PET data (Additional file 1: Fig. S3). Then we performed meta-analyses using METAL [33], of joint-cohorts with those that only had summary statistics (Memento, MCSA, WRAP, BACS, Korean study, and MISSION-AD) to obtain the race-specific meta-analyses. To conduct the multi-ethnic meta-analysis, a standarderror (StdErr)-based approach was used from Metal [33]. The variant-level visualization was accomplished using LocusZoom [30] tool.

As the amyloid PET imaging was calculated using different traces and pipelines, the joint analyses could lead to false positive or negative findings. Similarly, because of the different distribution of the amyloid PET data, the Z-scores could lead to spurious results. Existing studies suggest joint analysis can be more powerful than metaanalysis for low frequency variants where dataset is comprised of divergent samples [35, 36]. On the other hand, meta-analysis approaches do not require the complete original data as they can be used with summary statistics information while accounting for difference in population size, and they are less demanding computationally [37]. In order to confirm that the Z-score normalization and the joint analyses lead to robust results, we performed a GWAS analyses using raw centiloid values and the Z-scores for the Knight-ADRC and ADNI (Additional file 1: Fig. S4). A very strong correlation in p-values and effect size (R > 0.89) was found in these comparisons indicating that Z-scores will lead to the same signals as the raw amyloid data. In addition to the above validation, we performed a more systematic analyses comparing the *p*-values and effect sizes of the joint (z-scores) and the meta-analysis for all cohorts for those having raw phenotypic data available (A4=3,180, ADNI=1134, ADNIDOD=169, AIBL=1214, DIAN=209, HABS=258, Knight-ADRC=1,048, UPitt=345). Briefly, (1) the raw phenotype values were converted to log10 scale for each cohort and each cohort was analyzed independently. Then the cohort-specific results were combined by metaanalyses. For meta-analysis, we used METAL software and standard-error (SE)-based meta-analysis approach. (2) In parallel, the raw phenotypes were also converted to z-score, separately for each cohort, combined by performing a single joint analyses. (3) Finally, the *p*-values and effect sizes of associations were compared between these two analyses (joint vs. meta-analysis). We assessed the correlation of effect sizes and p-values at different *p*-values thresholds of associations (P=0.05,  $5 \times 10^{-05}$ , and  $5 \times 10^{-08}$ ) and found very strong correlation (R > 0.94) in all comparisons (Additional file 1: Fig. S5). This high correlation indicates that the joint and the meta-analyses lead to the same results.

We also performed *APOE*  $\varepsilon$ 4-, sex- and case-control status-stratified analyses. Most of these analyses were performed only on those datasets for which amyloid PET and genetic data were available. Therefore, the samples

sizes for those analyses were lower than for the overall analyses. We also compared the *p*-values of all these analyses using the z-scores joint vs the meta-analysis (Additional file 1: Fig. S6, S7 and Table S2), finding very high correlation.

In order to compare the effect size (BETA) across different ethnicities, we used two-sample t-test that takes into account the BETA and standard error (SE) for performing a pair-wise comparison and provide the *p*-value of significance whether difference between BETA is statistically significant or not. We developed a custom R function that uses BETA and square root of SE<sup>2</sup> of two groups to obtain a Z-score. As this Z-score follows a normal distribution with a sample size > 100 in each ancestry, we obtained a *p*-value for significance by using the pnorm function in R.

## **Post-GWAS** analyses

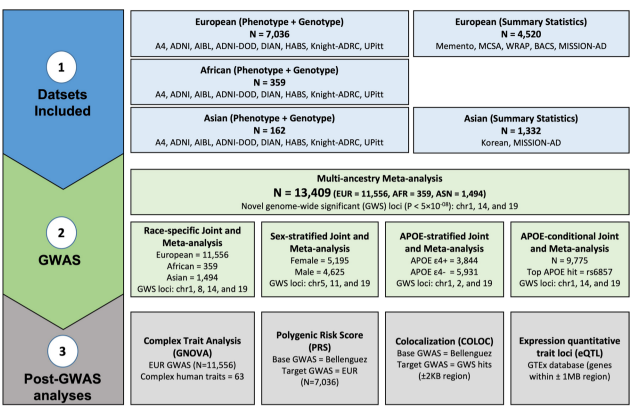
Multiple post-GWAS analyses were conducted for the functional annotation of the identified hits. A schematic overview of study design and conducted analyses is provided in Fig. 1.

#### Genetic covariance analysis

We used Genetic Covariance Analyzer (GNOVA) program [38] for assessing the genetic covariance and correlation of amyloid PET AD endophenotype with different complex human traits. Publicly accessible GWAS summary statistics data was downloaded for 63 different complex human traits. Specific details about the sample size and source of summary statistics for each trait are presented in Additional file 1: Table S3. The obtained summary statistics data was used to assess the genetic covariance of each trait with amyloid PET GWAS from non-Hispanic Whites (NHW) ancestry (N = 11,556). The tool (GNOVA) that has been used for calculating genetic correlation has the inherent ability to take the sample overlap between different GWAS into account. The algorithm allows random error terms  $\epsilon$  and  $\delta$  to be correlated in order to account for the non-genetic correlation introduced by sample overlapping between two GWAS.

## Bayesian co-localization analysis

We performed co-localization analysis using COLOC [39] tool (version 5.2.1). For each genomic locus of interest, identified by multi-ethnic or sex-stratified amyloid PET GWAS analyses, co-localization was assessed against AD risk GWAS from Bellenguez et al. [5]. Furthermore, to assess the genotype-specific expression of identified SNPs in human brain tissues, we performed cis-expression quantitative trait loci (QTL) analysis using gene expression data from the Genotype-Tissue Expression (GTEx) portal [40]. For



**Fig. 1** Schematic overview of datasets and performed analyses. Amyloid PET endophenotype and corresponding genotype data was available for 8 different cohorts with a total sample size of 7,557 (NHW = 7036, African = 359, Asian = 162). We also got GWAS summary statistics data from 6 external cohorts having a total sample size of 5852 (NHW = 4520, Asian = 1332). We performed Race-specific linear regression using amyloid PET as a quantitative endophenotype and age, sex, cohort name, and first ten genetic PCs as model covariates. The same analytic pipeline was used by the external cohorts for generating the summary statistics data. We meta-analyzed the results from internal and external summary statistics using a standard error (StdErr)-based meta-analysis approach using METAL software (N = 13,409). Furthermore, different post-GWAS analyses were carried out to identify novel SNPs associated with brain amyloidosis

both these analyses, posterior probability was calculated for five different hypotheses, (i) H0: there is no causal variant for either trait in the specified region, (ii) H1: there is a causal variant for trait 1, (iii) H2: there is a causal variant for trait 2, (iv) H3: both traits have distinct causal variants, and (v) H4: there is a single causal variant common to both traits. Here, our hypothesis of interest was H4. In case of co-localization with AD risk, a candidate locus was defined as 500 KB upstream and downstream of the variant of interest, however, in the case of eQTL analysis this region was extended to 1 MB on either side. For eQTL analysis, 10 types of tissues were considered from the GTEx portal, including "Brain Hypothalamus", "Brain Cerebellar Hemisphere", "Brain Cerebellum", "Brain Cortex", "Brain Amygdala", "Brain Caudate basal ganglia", "Brain Nucleus accumbens basal ganglia", "Brain Anterior cingulate cortex BA24", "Brain Putamen basal ganglia", and "Whole Blood".

## Polygenic risk score analysis

PRSice-2 software [41] was used for calculating the Polygenic risk score (PRS). We used the summary statistics data from the largest available clinically assessed AD case-control GWAS study [5] (N=788,989) as the base GWAS. We generated genetic risk scores as the weighted sum of the risk alleles for all participants in the amyloid PET GWAS of NHW ancestry (N=7036). The standard clumping and thresholding (C+T) approach was used where only those markers are retained that are most strongly associated with the disease. PRS calculation was generated at multiple different thresholds, ranging from  $5 \times 10^{-08}$  to 0.5 with an interval of  $5 \times 10^{-05}$ , on linkage disequilibrium (LD)-clumped SNPs by retaining the SNP with the smallest *p*-value and excluding the variants with r2>0.1 in a 250-kb window. We also calculated PRS by excluding the APOE region (GRCh38chr19:43,907,927-45,908,821) because of the high LD in this region.

## Results

#### Study design

In this study, we harmonized and integrated brain amyloid imaging and genetic data from 14 different cohorts with a total sample size of 13,409 (Table 1, Fig. 1), from three different populations (Non-Hispanic Whites (NHW), African (AFR) and Asian (ASN)), in order to identify genetic variants associated with amyloidosis and AD risk. The datasets with individual level genetic and brain amyloid imaging data and from the same ethnicity were analyzed using a joint approach after harmonizing the amyloid imaging. Results from these analyses were then meta-analyzed together with the datasets from the same ethnicity for which only summary statistics were available leading to race-specific brain amyloid imaging GWAS. Multi-ethnic GWAS was then performed using a random effect model. Moreover, APOE-, sex-, and diagnosis-stratified analyses were performed including those datasets with individual-level data to identify additional loci and genes to further explore the genetic architecture of brain amyloidosis. Functional annotation and expression and protein Quantitative Trait Loci (pQTL) mapping was performed to identify the most likely functional genes driving the associations for the GWAS signals. Finally, the overlap between the genetic architecture of brain amyloid imaging with AD risk and other traits were analyzed using Polygenic Risk Scores (PRS) and genetic covariance analysis.

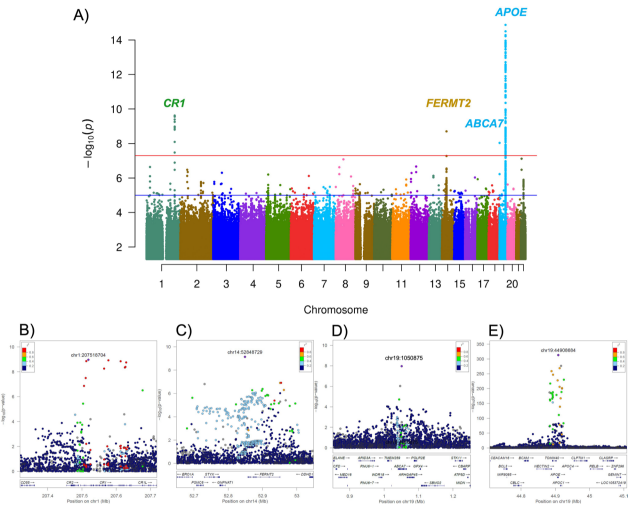
## **GWAS** analyses

The race-specific meta-analysis for NWH participants (n=11,556) from 13 different cohorts yielded a very strong signal in the APOE locus. In this locus, rs429358, which codifies for APOE ɛ4, was the most significant SNP  $(P=1.8\times10^{-416}, \beta=0.62, SE=0.01, MAF=0.19, I^2=99.7;$ Additional file 1: Fig. S8A). Besides the APOE locus, we identified two additional novel genome-wide significant signals in chr14q.22.1 (rs117834516,  $P = 8.7 \times 10^{-09}$ ,  $\beta = 0.15$ , SE = 0.03, MAF = 0.06, I<sup>2</sup> = 0) on the *FERMT2 locus* and chr1q.32.2 (rs6656401, P=9.1×10<sup>-10</sup>,  $\beta$ =0.10, SE = 0.02, MAF = 0.18, I<sup>2</sup> = 43.1) on the CR1 locus (Additional file 1: Fig. S9C, D). All these signals colocalized with that of AD risk GWAS [4] having posterior probability of 99% for sharing a single causal variant (Additional file 1: Table S4). Colocalization with eQTL (in multiple brain regions based on GTEx [42]) was also observed for the chr1q.32.2 locus, rs6656401, with CR1 mRNA levels (PP.H4=0.99), the chr14q.22.1, rs117834516, with STYX mRNA levels (PP.H4=0.93), but not FERMT2 (PP. H4=0.08) (Additional file 1: Table S5, S6). Recent GWAS have provided strong evidence for CR1 and FERMT2 being the risk factors for the development of AD [43, 44]. Specifically, *CR1* has been reported to be consistently involved in immune system related pathways, particularly complement and inflammatory cytokines [45]. Changes in the gene expression level of *FERMT2* in the neurons have been shown to effect both extracellular A $\beta$ 40 and A $\beta$ 42 as well as phospho-Tau [46].

In case of AFR-specific analyses (n=359), rs429358 was found to be the most significant SNP in the APOE locus (P= $1.0 \times 10^{-11}$ ,  $\beta = 0.49$ , SE=0.07, MAF=0.21; Additional file 1: Fig. S8B). In addition, we found a novel locus on chr8q.22.1 (rs2271774, P= $2.5 \times 10^{-09}$ ,  $\beta = 1.10$ , SE = 0.17, MAF = 0.028) that passed the genome-wide significance. This same SNP has a MAF of 1.3% in ASN  $(N = 336, P = 0.84, \beta = -0.04, SE = 0.19, MAF = 0.013)$  and 0.21% in NHW (N=6329, P=0.84,  $\beta$ =0.13, SE=0.56, MAF = 0.0021) but it was not even nominally associated in these two ancestries, suggesting it to be an AFR-specific signal, or a false positive. Additional studies having larger dataset, focused on individuals with AFR background, will be needed to further replicate this association. This SNP is located in the PTDSS1 gene-region that also includes GDF6, UQCRB, MTERF3, and SDC2 genes (Additional file 1: Fig. S10A). Although we did not find any association of chr8q.22.1 with gene or protein levels QTLs (Additional file 1: Table S7), we found that PTDSS1, but not any other gene in this region, to be significantly differentially expressed in different brain tissues (IFG, PHG, STG, and TCX) with a consistent negative log2 fold change [47], nominating PTDSS1 as the functional gene in this locus.

In case of ASN ancestry-specific analyses (n=1,494), no additional genome-wide signals were found besides *APOE* locus (P=6.7×10<sup>-28</sup>,  $\beta$ =0.14, SE=0.01, MAF=0.14; Additional file 1: Fig. S8C). Overall, the rs429358 SNP that codifies for *APOE*  $\epsilon$ 4 was the most significant SNP across every race-specific GWAS with a consistent positive effect size.

The multi-ethnic meta-analysis of NHW, AFR, and ASN ethnicities across 14 different cohorts (n = 13,409)further validated the results from individual ancestry GWAS and revealed one additional locus passing the genome-wide significance threshold on chr19p.13.3 (rs12151021,  $P = 9.2 \times 10^{-09}$ ,  $\beta = 0.07$ , SE = 0.01, MAF = 0.32,  $I^2$  = 0; Fig. 2) located near the *ABCA7* gene (Additional file 1: Fig. S9B). This same SNP is strongly associated with AD risk (P= $4.1 \times 10^{-30}$ ,  $\beta = 0.1$ ), and colocalizes (PP.H4=0.99) with AD risk [4]. This gene has already been shown to affect cognitive and behavioral aspects of AD [48-50]. We did not observe colocalization of this signal with eQTL for ABCA7 mRNA levels (PP.H4 < 0.05) based on GTEx [42], however, nearby genes such as RNU6-2 and GRIN3B were



**Fig. 2** Multi-ethnic meta-analysis (N = 13,409) identified novel signals in chr 1, 14, and 19 associated with brain amyloidosis. **A** Manhattan plot showing the *p*-values in the multi-ethnic meta-analysis. The blue and red lines represent the suggestive (P =  $1 \times 10^{-5}$ ) and genome-wide significance thresholds (P =  $5 \times 10^{-8}$ ). Variants with a *p* value below  $1 \times 10^{-15}$  are not shown. Local Manhattan plot for the chr1 (**B**), chr14 (**C**), and chr19 (**D**, **E**) loci. The relative location of genes and the direction of transcription are shown in the lower portion of the locus zoom plots

colocalized with rs12151021 in brain frontal cortex (PP. H4=0.93) and brain cortex (PP.H4=0.94), respectively (Additional file 1: Table S8).

#### APOE conditional analyses

The *APOE* locus that encodes apolipoprotein E (*APOE*) is a strong genetic risk factor for AD. Previous studies have shown that *APOE*  $\varepsilon$ 4 individuals have higher amyloid burden [51, 52]. Contrarily, *APOE*  $\varepsilon$ 2 carriers, showing reduced A $\beta$  deposition [53], have been associated with milder clinical and pathological AD when compared to  $\varepsilon$ 4 carriers [54].

In order to determine if the signals observed in the current analysis are driven by *APOE*  $\varepsilon 2/\varepsilon 3/\varepsilon 4$ , we performed

*APOE* conditional analyses using the summary statistics data from the multi-ethnic meta-analysis (N=13,409), using COJO [55]. In the multi-ethnic meta-analysis, rs429358, which codifies *APOE*  $\varepsilon$ 4, was found to be the most significant SNP from chr19 (P= $6.2 \times 10^{-311}$ ,  $\beta$ =0.35, SE=0.01, MAF=0.19). Therefore, a conditional analysis was performed by adjusting for rs429358 (*APOE*  $\varepsilon$ 4) and rs7412 (*APOE*  $\varepsilon$ 2). A total of 50 SNPs on the *APOE* locus remained genome-wide significant at P<5×10<sup>-08</sup> (Additional file 1: Fig. S11 and Table S9), and the most significant was rs73052335 (Additional file 1: Fig. S12A). After conditioning for *APOE*  $\varepsilon$ 4, *APOE*  $\varepsilon$ 2, and rs73052335, we identified three additional independent signals in this region: (rs1081105/chr19:44,909,698:A:C; rs438811/chr19:44,913,484:C:T;

rs4420638/chr19:44,919,689:A:G) that remained genome-wide significant in the conditional analyses, indicating that there are up to six independent signals in the APOE locus (Additional file 1: Table S9 and Fig. S12). We performed functional annotation and expressionand splicing-QTL mapping for these independent signals and several of these SNPs are either eQTL or sQTL for TOMM40 and NECTIN2, suggesting that these genes may also be contributing to amyloid deposition independently of APOE. Colocalization analysis with AD risk for all the hits from APOE conditional analyses (N=50)with AD risk GWAS [4] suggests more than half of these signals to be colocalizing with H4>0.8 (Additional file 1: Table S10).

We also performed similar analyses including only the datasets from which we have individual level data (N=7,557), but instead of conditioning on the SNPs that codified for *APOE*  $\epsilon$ 4, *APOE*  $\epsilon$ 2, we conditioned on the full *APOE* genotypes:  $\epsilon$ 22,  $\epsilon$ 33,  $\epsilon$ 24,  $\epsilon$ 34 and  $\epsilon$ 44, codified as 0, 1, 2, 3, 4 and 5 respectively (Additional file 1: Fig. S13). One independent and additional signal (sentinel SNP: rs5117/chr19:44,915,533:T:C) passed genomewide significance (P=1.6×10<sup>-08</sup>,  $\beta$ =0.11, SE=0.02, MAF=0.29), after condition for full *APOE* genotype. This SNP is in high LD with rs73052335 (D'=1, R<sup>2</sup>=0.23, Additional file 1: Table S11) and likely to correspond to the same signal.

#### Race-specific APOE analyses

In order to assess whether *APOE* exhibits a race-specific effect, we analyzed the association between quantitative amyloid PET levels and *APOE*-associated variants on chr19q.13.32 (*APOE*  $\epsilon$ 4/rs429358, *APOE*  $\epsilon$ 2/rs7412, and rs5117). *APOE*  $\epsilon$ 4 SNP showed a consistent and positive association with amyloid PET levels (positive effect size in each of the analyzed races; NHW=0.62, AFR=0.49, and ASN=0.14; Additional file 1: Table S12). The effect size on *APOE*  $\epsilon$ 4 in ASN was significantly lower than the NHW (two sample *t*-test, P=3.8×10<sup>-242</sup>) and AFR (two sample *t*-test, P=7.4×10<sup>-07</sup>) ethnicities.

We observed a protective role of *APOE*  $\varepsilon$ 2 variant across all ethnicities with a consistent negative effect size (NHW = -0.33, AFR = -0.16, ASN = -0.02; Additional file 1: Table S12). Although the difference in the effect size was only significant in NHW and ASN ethnicities (two sample *t*-test, P=8.1×10<sup>-18</sup>), AFR had almost half the effect size of NHW, indicating a race-specific effect of *APOE*  $\varepsilon$ 2.

The third independent signal in the *APOE* locus, rs5117, consistently positively associated with brain amyloidosis (NHW=0.41, AFR=0.04, and ASN=0.11; Additional file 1: Table S12), also exhibited a race-specific effect. The observed effect size was significantly

higher in NHW as compared to AFR (two sample *t*-test,  $P=3.3 \times 10^{-09}$ ) and ASN (two sample t-test,  $P=1.9 \times 10^{-93}$ ) ethnicities. The fourth independent signal in the *APOE* locus, rs73052335, also showed consistently positive association across all races (NHW=0.56, AFR=0.45, and ASN=0.43), but the observed effect size was not race-specific. Overall, this data indicates that *APOE* is a complex locus in which multiple race-specific signals influence brain amyloidosis.

### APOE ε4 stratified analyses

Considering the well-established risk effect of *APOE*  $\varepsilon 4$ and protective effect of *APOE*  $\varepsilon 2$  alleles in AD, we asked if amyloid positivity also shows any *APOE*  $\varepsilon 4$ -dependent difference. Therefore, we performed an *APOE*  $\varepsilon 4$ -stratified analysis and observed genome-wide significant SNPs in both *APOE*  $\varepsilon 4$ + (N=3,844) and *APOE*  $\varepsilon 4$ -(N=5,931) strata (Additional file 1: Fig. S14).

In the *APOE*  $\epsilon$ 4+ strata, the top hit on the *APOE* locus was driven by the *APOE*  $\epsilon$ 4 allele (rs429358,  $P=1.5 \times 10^{-26}$ ,  $\beta=0.37$ , SE=0.03, MAF=0.19), which is capturing the association of one vs two *APOE*  $\epsilon$ 4 alleles. The signal in the *CR1* locus, chr1q.32.2 (rs679515,  $P=2.7 \times 10^{-08}$ ,  $\beta=0.15$ , SE=0.03, MAF=0.18) was in high LD (R<sup>2</sup>=0.97) with previously observed signal in the NHW and multi-ethnic meta-analysis (chr1q.32.2), also remained significant in the *APOE*  $\epsilon$ 4+ strata, but only nominally significant in the *APOE*  $\epsilon$ 4- (P=0.02,  $\beta$ =0.05, SE=0.02). We found that this SNP also has a significant interaction with *APOE*  $\epsilon$ 4+ (P=0.008) and the effect size in the *APOE*  $\epsilon$ 4+ vs  $\epsilon$ 4- is also significantly different (p=0.001; Additional file 1: Table S13), indicating an interaction with *APOE*.

In case of the *APOE* e4- strata, the *ABCA7* locus (chr19p.13.3) passed the genome-wide significance threshold (rs12151021, P=3.3×10<sup>-09</sup>,  $\beta$ =0.10, SE=0.02, MAF=0.18; Additional file 1: Fig. S14A), but was only nominal significant in the *APOE* e4+ strata (P=0.02,  $\beta$ =0.05, SE=0.02). We did not find any significant interaction between this SNP and *APOE* e4, nor was the effect size in e4+ vs *APOE* e4- strata significantly different (Additional file 1: Table S13). Similar findings were identified for *FERMT2*, with a more significant *p*-value in *APOE* e4- (P=7.3×10<sup>-06</sup>,  $\beta$ =0.16, SE=0.03) compared to *APOE* e4+ (P=0.02,  $\beta$ =0.11, SE=0.05) but without interaction with *APOE* e4.

Notably, in the *APOE*  $\epsilon$ 4- strata we identified rs1065853 as the most significant SNP on the *APOE* locus (P=1.5×10<sup>-13</sup>,  $\beta$ =-0.21, SE=0.03, MAF=0.07). This SNP is in high LD with *APOE*  $\epsilon$ 2 (D'=0.99, R<sup>2</sup>=0.96). In addition, we identified a novel signal on chr2q.12.2 (rs567226423; P=2.8×10<sup>-08</sup>,  $\beta$ =0.81, SE=0.15, MAF=0.003). Functional annotation by VEP

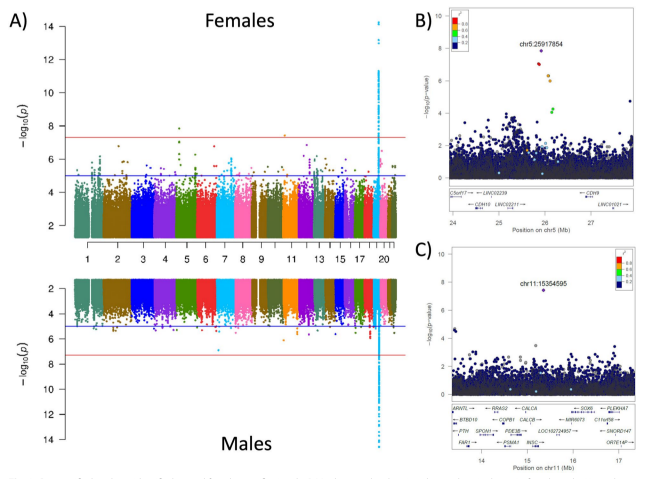
[56] suggest this SNP to be a regulatory region variant for *UXS1* protein coding gene, however, another nominally significant SNP, chr2q.12.2 (rs191708024, P=1.9×10<sup>-04</sup>,  $\beta$ =0.28, SE=0.08, MAF=0.01) within high LD (D'=1, R<sup>2</sup>=0.50), is an eQTL for *NCK2, which is* a known AD risk gene [5]. Additional studies even in larger dataset are needed to replicate this finding.

## Sex-stratified GWAS

About two thirds of people diagnosed with AD are women, however, the life expectancy for women is longer than for men [57]. Furthermore, age is the greatest risk factor for AD dementia as chances for developing AD by the age of 45 is 20% and 10% for women and men, respectively [57]. In light of these facts, we wanted to assess if amyloid burden also exhibits sex-dependent differences

by performing the association analysis separately for the female (N = 5195) and male (N = 4625) strata.

No genome-wide signals were found in males, but the female-specific analyses revealed two independent genome-wide significant SNPs on chr5p.14.1 (rs529007143,  $\beta = 0.79$ , SE=0.14, P=1.4×10<sup>-08</sup>, sexinteraction  $P = 9.8 \times 10^{-07}$ , MAF = 0.006,  $I^2 = 0$ ; Fig. 3) and chr11p.15.2 (rs192346166,  $\beta = 0.94$ , SE = 0.17,  $P = 3.7 \times 10^{-08}$  $P = 1.3 \times 10^{-03}$ , sex-interaction MAF=0.004,  $I^2=0$ ). We looked for SNPs in high LD with these signals and found several SNPs in high LD that were also highly significant, indicating them to be potential new signals and not an imputation error or artifact (Additional file 1: Table S14). In addition, in the sex-AD interaction analysis, both of the identified variants passed the suggestive significance (Additional file 1: Fig. S15) and showed a positive association with



**Fig. 3** Sex stratified analyses identified several female specific signals. **A** Manhattan plot showing the *p*-values in the 5195 female and 4625 male participants across 9 cohorts. The blue and red lines represent the suggestive ( $P = 1 \times 10^{-5}$ ) and genome-wide significance thresholds ( $P = 5 \times 10^{-8}$ ). Variants with a p value below  $1 \times 10^{-15}$  are not shown. The observed genomic control value ( $\lambda$ ) was 1 for both strata. Local Manhattan plot showing the genome-wide significant locus from the chr5 (B) and chr11 (C) for female-specific signals. The relative location of genes and the direction of transcription are shown in the lower portion of the locus zoom plots

amyloid deposition for rs529007143 ( $\beta$ =0.54, SE=0.11,  $P = 9.8 \times 10^{-07}$ ) and chr11p.15.2 (rs192346166,  $\beta = 0.41$ , SE=0.13, P= $1.3 \times 10^{-03}$ ) variants. Functional annotation of these rare genetic variants indicates that the intergenic upstream genes are MSNP1 and INSC/EHF for rs529007143 and rs192346166, respectively (Additional file 1: Fig. S16). Expression and pQTL analyses were not informative to identify the most likely functional genes in those regions (Additional file 1: Table S15). We also did not find any overall gene-expression change between AD cases and controls for MSNP1. However, gene expression of *EHF* is higher in AD brains compared to controls, and more interestingly this difference is also driven by women [47]. The identified loci did not colocalized with AD risk GWAS (Additional file 1: Table S4), probably because the considered AD risk GWAS has both males and females individuals. To conclude, we have identified two novel female-specific signals that are positively associated with amyloid deposition, however, additional studies with even a larger sample size are needed to replicate these finding and proper colocalization analyses are also needed in female-only GWAS for AD risk to further validate these signals.

## Case-control stratified analyses

Population-based case–control studies have become increasingly popular for finding common polymorphisms that underlie complex human traits. In order to identify such genetic variants underlying our AD endophenotype of interest (Amyloid PET), we conducted a case–control stratified association analysis (CO=5846, AD=1138) and identified multiple variants from the *APOE* locus that passed the genome-wide significance in both strata (Additional file 1: Fig. S17).

Briefly, in these analyses only the APOE locus passed the genome-wide significance, with APOE £4 (rs429358) being the most significant SNP. Even, the effect size in cases and controls (AD:  $\beta = 0.59$ , SE = 0.02,  $P = 1.2 \times 10^{-165}$ , MAF = 0.19; CO:  $\beta = 0.47$ , SE = 0.04,  $P=1.9\times10^{-29}$ , MAF=0.23) were in the same direction, the statistical analyses indicate that they are significantly different (p=0.007, Additional file 1: Table S16). Similar findings were observed for the protective effect of APOE  $\epsilon 2$  (AD:  $\beta = -0.54$ , SE=0.1, P=3.7×10<sup>-08</sup>; CO;  $\beta = -0.26$ , SE = 0.03, P = 2.1 × 10<sup>-16</sup>), with cases showing significantly higher effect size than controls (p=0.007; Additional file 1: Table S16). We observed similar and non-significantly different effect sizes for CR1, FERMT2, and *ABCA7* in AD and CO (p > 0.29), indicating that both strata contribute to the association.

We also checked the overlap between known AD risk loci [5] and the case-control stratified GWAS. In the case-only analysis (N=1138), we found seven

SNPs to be nominally significant with consistent effect size direction except for one variant (Additional file 1: Table S17). The most significant was on chr14q.32.12/rs7401792 (*SLC24A4*, P=0.0010,  $\beta$ = -0.14, SE=0.04, MAF=0.37; Additional file 1: Table S17). In case of controls (N=5846), we identified 11 SNPs to be nominally significant, with chr7p.21.3 (rs6943429) being the most significant (*UMAD1*, *p*=0.0010,  $\beta$ =0.05, SE=0.02, MAF=0.42; Additional file 1: Table S17). Overall, we found more loci to be associated with AD risk in the amyloid PET control GWAS.

## Association of known AD risk loci with amyloid burden

Next, we assessed if there is an overlap between the genetic architecture of amyloid imaging and AD risk beyond the identified SNPs in the APOE, ABCA7, FERMT2, and CR1 loci that all colocalized with AD risk (PP.H4>0.88; Additional file 1: Table S4). Among the 82 genome-wide significant sentinel SNPs associated with AD risk reported by Bellenguez et al. [5], 18 additional AD risk variants were also nominally associated (P < 0.05) with brain amyloidosis in the current amyloid PET GWAS (Additional file 1: Table S18). This overlap of overall 21 SNPs between AD risk and amyloid PET GWAS was found to be statistically significant (hypergeometric test  $P = 4.1 \times 10^{-12}$ ). The identified SNPs included key AD associated genes, such as ANK3, APP, BIN1, CLU, FERMT2, and TREM2. Importantly, for most of the identified SNPs, the direction of effect size ( $\beta$ ) was consistent between Bellenguez et al. [5] and the current amyloid PET GWAS. These results suggest that amyloid imaging endophenotype can serve as a proxy for AD risk.

#### Polygenic risk score analysis

In order to further determine the association of brain amyloidosis and AD risk, we determined if AD Polygenic risk score (PRS) were associated with amyloid PET. PRS were calculated with and without APOE region (GRCh38, chr19:43,907,927-45,908,821) from the most recent AD case-control GWAS [5]. We observed significant association between PRS for AD risk and brain amyloidosis (PRS.R2=0.05, P= $2.6 \times 10^{-85}$ ) at P-value threshold (P<sub>T</sub>) equal to genome-wide significance  $(P_T = 5 \times 10^{-08};$  Additional file 1: Fig. S18) when APOE was included. We also observed a significant association (PRS.R2=0.01, P= $1.4 \times 10^{-19}$ ) even when the APOE region was removed from the reference GWAS. Overall, age, sex, cohort, first ten genetic principal components, and the polygenic score explained 8.7% (Full.R2=0.087) and 4.7% (Full.R2=0.047) of the variance in amyloid levels with and without APOE, respectively. These results further strengthen our hypothesis that there are loci besides APOE that are associated with amyloid burden in

the brain and they also contribute to the observed association with AD risk PRS.

## Genetic covariance between amyloid PET GWAS and complex human traits

To assess whether amyloid PET endophenotype shares its genetic basis with other complex human traits, we performed genetic covariance analyses using summary statistics from amyloid PET GWAS from NHW ancestry (N=11,556) and 63 human health-related phenotypes [58] (Additional file 1: Table S3). Pair-wise genetic covariances between amyloid PET AD endophenotype and 63 human health-related phenotypes are shown in Fig. 4 and Additional file 1: Table S19. In total, 7 phenotypes showed significant correlation after multiple testing and sample overlap correction.

We observed a strongly positive correlation with high-density lipoprotein (HDL; cor=0.29, SE=0.09,  $FDR = 9.2 \times 10^{-05}$ ), Total cholesterol (cor = 0.19,SE = 0.09,  $FDR = 2.5 \times 10^{-02}$ ), and Asthma (cor = 0.15, SE=0.06, FDR= $2.5 \times 10^{-02}$ ), among others. Moreover, we observed significantly negative correlations with Epilepsy (cor = -0.38, SE = 0.13, FDR =  $2.5 \times 10^{-03}$ ) and Celiac disease (cor = -0.16, SE = 0.06, FDR =  $2.2 \times 10^{-02}$ ). Removing the APOE region had minimal effect on the pair-wise genetic correlations and their significance, with the exception of identifying a new significant correlation for Multiple Sclerosis (cor = -0.23, SE = 0.06,  $FDR = 7.3 \times 10^{-03}$ ). Taken together, these results provide evidence of consistency in the observed polygenic trends across comparable measures from independent datasets.

## Discussion

In this study we explored the genetic basis of brain amyloidosis by analyzing the largest collection of amyloid PET imaging data to date. We used harmonized amyloid PET levels as a quantitative trait to conduct the racespecific GWAS across eight different cohorts (N=7557). Further, the summary statistics data from 6 additional cohorts were included to perform multi-ethnic metaanalysis for identifying novel genetic associations with brain amyloidosis (N=13,409).

The meta-analysis revealed a very strong significant genome-wide locus on the *APOE* gene region, validating previous findings [11, 14, 15]. We leveraged these large and multi-ethnic cohorts to further disentangle the role of *APOE* in brain amyloidosis and AD risk. It is known, and well accepted that there are at least two independent signals in the *APOE* gene:  $\varepsilon$ 4 and  $\varepsilon$ 2 [11, 15]. We found a strong and positive association of  $\varepsilon$ 4 across populations (Effect size NHW=0.62 [SE=0.01], AFR=0.49 [SE=0.07], ASN=0.14 [SE=0.01]; Additional file 1: Table S12), even though the MAF and the effect size

were significantly different across populations. We also observed significant heterogeneity in the effect of *APOE*  $\epsilon_2$ , with strong protective effect of this variant in NHW (Effect size=-0.33, SE=0.02), but almost nothing at all in ASN (Effect size=-0.02, SE=0.03), and AFR showing an intermediate but not significant association of this variant with brain amyloidosis (Effect size=-0.16, SE=0.10). As our population principal component factor analyses indicate that our AFR population includes samples with some NHW contribution (Additional file 1: Fig. S1B), it will be important to perform future analyses evaluating local admixture mapping to determine the effect size of the *APOE*  $\epsilon_2$  in individuals from a NHW or AFR background.

In addition to the known ɛ4 and ɛ2 independent variants, our APOE conditional analyses demonstrate that there are at least four additional risk signal in this locus on chr19q.13.32, tagged by rs73052335 ( $\beta = 0.28$ , SE = 0.02, P =  $2.1 \times 10^{-71}$ , MAF = 0.17)/rs5117 ( $\beta$  = 0.07,  $P = 1.2 \times 10^{-13}$ , MAF = 0.29), rs1081105 SE = 0.01,  $P = 6.4 \times 10^{-24}$ , MAF = 0.04), $(\beta = 0.29,$ SE = 0.03, SE = 0.01,  $P = 1.6 \times 10^{-57}$ , rs438811  $(\beta = -0.15,$ MAF = 0.41), and rs4420638 ( $\beta$  = -0.06, SE = 0.001,  $P = 1.0 \times 10^{-09}$ , MAF = 0.23), that have also been reported to be associated with AD risk with positive colocalization (Additional file 1: Table S10). Similar to APOE ɛ2, rs5117 signal shows very specific racespecific effects. This variant has a strong effect in NWH, a significant but weaker effect on ASN and no effect on AFR. Based on our analyses, it is not clear if this variant is modifying amyloid imaging through an APOE-dependent mechanism or by some other gene. Our eQTL analyses indicate that the additional APOE variants also regulate TOMM40 or NECTIN2 mRNA levels. TOMM40 (translocase of outer mitochondrial membrane 40 homolog) has also been reported to be associated with LOAD and multiple LOAD-related neuroimaging phenotypes in the 19q13 region [59-61]. A poly-T polymorphism in TOMM40 has been proposed to modify AD risk independently of APOE in multiple studies, but with contradictory results [59-62]. Nectin cell adhesion molecule 2 (NECTIN2), an important mediator of immune system, has already been shown to be downregulated in the neurons of AD cases [63] and different genetic variants in this gene have been associated with alterations in the CSF A $\beta$  and tau levels as well as AD [2, 64, 65]. In addition, common variants in NECTIN2, have shown a strong association with CSF sTREM2 [66]. Besides APOE ɛ4 and ɛ2, various other variants have also been shown to exhibit protective or risk effect on AD. For example, a recent report suggest that the individuals having two copies of APOE e3 Christchurch (p.R136S) mutation may slow down the

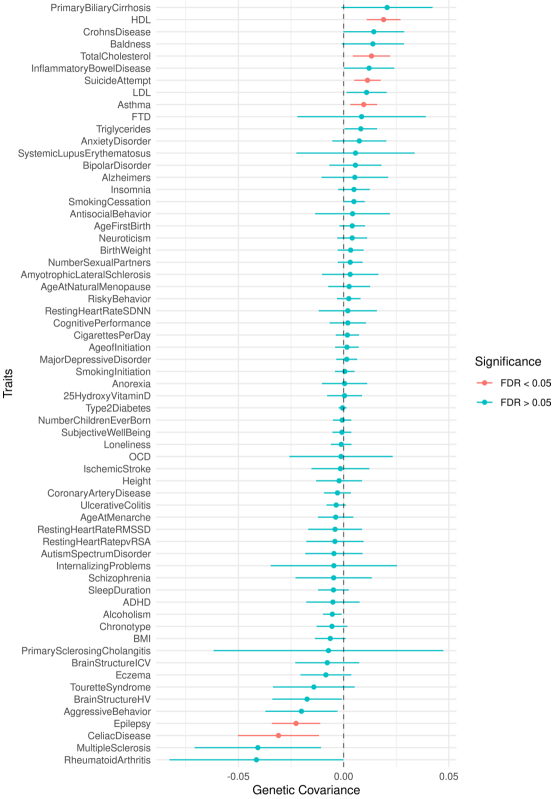


Fig. 4 Genome-wide genetic covariance results. Genetic covariance between multi-ethnic amyloid PET GWAS (NHW = 11, 816) and 63 complex human traits. Error bars represent 95% confidence intervals

progression of AD by preventing the accumulation of tau tangles and associated cell death [67]. Furthermore, multiple signals have been found in the *APOE* locus for CSF  $A\beta_{1-42}$  after controlling for *APOE* genotype and adjusting for multiple comparisons based on Bonferroni threshold [68].

The effect of APOE ɛ4 and ɛ2 with AD risk and AD endophenotypes is so strong that identifying, validating, and characterizing additional independent signals in this region remains challenging and additional studies are needed. In any case, these results strength the notion that multiple independent risk and protective variants are found in the APOE locus, and more importantly that these signals show a very clear race-specific effect. Other studies have also reported a race-specific effect of APOE on AD risk [69], mainly for APOE e4. Here we demonstrate that this observation can be extended to amyloid imaging but more importantly, we demonstrate that this differential effect is more pronounced to APOE e2 and for the new additional signal we identified. This has important implications when estimating the individual risk at a population level that may affect future clinical trial strategies and therapies. The new results from the Lecanemab study [70] indicates that this treatment has different effects depending on the ethnicity and APOE genotype. The results from this study could help to further understand the result from this trial.

Besides identifying multiple independent signals in the APOE locus, we also found two additional NHW signals in the CR1 (chr1q.32.2,  $P=9.1 \times 10^{-10}$ ,  $\beta=0.1$ , SE=0.02, MAF=0.18) and FERMT2 (chr14q.22.1,  $P = 8.7 \times 10^{-09}$ ,  $\beta = 0.15$ , SE = 0.03, MAF = 0.06) loci, and a multi-ethnic signal in ABCA7 (chr19p.13.3,  $P = 9.2 \times 10^{-09}$ ,  $\beta = 0.07$ , SE = 0.01, MAF = 0.32) locus. All these signals have already been shown to be significantly associated  $(CR1 = 5.2 \times 10^{-33}, FERMT2 = 5.8 \times 10^{-10},$  $ABCA7 = 4.1 \times 10^{-30}$ ) with AD risk [5] and our colocalization analyses indicate they all share a single causal variant (PP.H4>0.88; Additional file 1: Table S4). We observed a significant interaction between APOE and CR1 signal  $(P=0.008, \beta=0.05, SE=0.02;$  Tale S16). This signal was mainly driven by APOE  $\varepsilon 4$ +stratum (P=3.3×10<sup>-08</sup>,  $\beta = 0.15$ , SE = 0.03) but also nominally significant in the APOE  $\epsilon$ 4-stratum (P=0.05,  $\beta$ =0.04, SE=0.02), and exhibited significant difference in the effect size between both these strata (two-sample t-test p = 0.001). We did not observe an interaction of APOE with FERMT2 and ABCA7 signal (p > 0.20), also their effect sizes were not significantly different (p > 0.10) between  $\epsilon 4 +$  and ε4- strata.

We found a strong overlap in the genetic architecture of AD risk and brain amyloid imaging. Alzheimer disease PRS with  $(P=2.6\times10^{-85})$  and without  $(P=1.4\times10^{-19})$ APOE showed a strong association with amyloid imaging. Additionally, 21 of the sentinel SNPs for the latest GWAS for AD risk [5] showed at least a nominal significance (p < 0.05) for brain amyloidosis. For most of the overlapping SNPs, the effect size directions were consistent across both analyses. Some of the most important genes that have been associated with these SNPs included, CR1 (Complement Receptor Type 1), ABCA7 (ATP-Binding Cassette Sub-Family A Member 7), BIN1 (Bridging Integrator 1), ANK3 (Ankyrin 3), TREM2 (Triggering Receptor Expressed On Myeloid Cells 2), and CLU (Clusterin). All these genes have been shown to be implicated in neuropathology and AD [40, 71–77]. For example, ABCA7 has been shown to mediate the generation of high-density lipoprotein (HDL) with apolipoproteins [78]. Like APOE, CLU is also known as a component of lipoprotein and implicated in a wide range of biological functions including cholesterol and lipid transport [79], however, the specific roles of these genes in brain cholesterol homeostasis and its involvement in AD is still unknown. Taken together, these results indicate that genetic architecture of amyloid PET endophenotype is very similar to that of AD risk, therefore, it may be used as a proxy for explaining the yet to be discovered genetic variance in AD.

Besides the robust association of brain amyloidosis with *APOE* locus, SNPs within *RBFOX1* gene on chr16p.13.3 have also been shown to reach genome-wide significance in previous studies (top SNP rs56081887,  $P=3\times10^{-09}$ ,  $\beta=0.61$ , MAF=0.09) [15]. We did not find a significant association for this variant in our race-specific GWAS for NHW ancestry (p=0.75,  $\beta=-0.01$ , SE=0.03; Additional file 1: Fig. S19) or multi-ethnic meta-analysis (p=0.67,  $\beta=-0.01$ , SE=0.03). As the initial association of this SNP was identified on controls only, we analyzed if this SNP showed an association in controls-only (N=5,846). We found rs56081887 (p=0.04,  $\beta=0.060$ , SE=0.03) and rs34860942 (p=0.03,  $\beta=0.064$ , SE=0.03) to be nominally associated with brain amyloidosis.

Additional interactions were found on our sex-specific analyses. Two new female-specific loci were detected in the sex-stratified multi-ethnic GWAS: MSNP1 gene on chr5.p14.1 (rs529007143,  $\beta = 0.79$ ,  $P = 1.5 \times 10^{-08}$ , MAF = 0.006) and EHF gene on chr11.p15.2 (rs192346166,  $\beta = 0.94$ ,  $P = 3.9 \times 10^{-08}$ , MAF = 0.004). Our analyses indicate that expression level of this gene is altered in AD brains, specifically in women, supporting the race-specific association. EHF (ETS Homologous Factor) is a DNA-binding transcription factor activity and RNA polymerase II cis-regulatory region sequencespecific DNA binding protein that play a crucial role in regulating epithelial cell differentiation and proliferation [80], however the role of this protein on AD risk is still unclear. Future studies, focused on sex-stratified analyses in AD, will be needed to replicate these findings and identifying the functional signals on chr11 and 5 to better understand the role of *EHF* on AD risk in females.

Another novel signal identified in this study is the AFR-specific signal on chr8q.22.1 (rs2271774,  $\beta = 0.98$ ,  $P = 7.8 \times 10^{-09}$ , MAF = 0.028) that, based on our functional mapping, is driven by PTDSS1. This gene encodes the enzyme phosphatidylserine synthase 1 (PSS1), which is involved in the transport of phospholipids between endoplasmic reticulum and mitochondria [81], and which is highly expressed in brain tissue, especially in neurons. This same SNP has a MAF of 1.3% in ASN  $(N=336, p=0.84, \beta=-0.04, SE=0.19, MAF=0.013)$  and 0.21% in NHW (N=6329, p=0.84,  $\beta=0.13$ , SE=0.56, MAF = 0.0021) but it was not even nominally associated in these two ancestries, suggesting it to be an AFR-specific signal or be a false-positive signal. Additional studies with an even larger AFR-specific sample size will be needed to further confirm this finding.

Besides the positive genetic overlap of amyloid PET AD endophenotype with different neurological disorders such as AD, Amyotrophic Lateral Sclerosis (ALS), and Frontotemporal dementia (FTD), we also found a significant genetic correlation with a number of different human traits. We observed strong genetic correlations with HDL and total cholesterol levels (Fig. 4 and Additional file 1: Table S19). It is also interesting that we observed significant negative correlations between AB deposition and brain structure hippocampus and intracranial volumes, validating existing findings that suggest brain volume loss during the mild cognitive impairment (MCI) to AD transition [82]. Notably, we observed strong positive genetic correlation between our trait of interest and different inflammatory disorders such as Crohn's disease (cor=0.07, SE=0.04, FDR < 0.19), ALS (cor = 0.03, SE = 0.06, FDR < 0.8), and Primary Biliary Cirrhosis (cor = 0.08, SE = 0.05, FDR < 0.21). Moreover, some genetic traits that were significantly negatively correlated were also related to the dysregulation of immune response e.g. Celiac disease (cor = -0.16, SE = 0.06,  $FDR < 2.2 \times 10^{-02}$ ) and Multiple Sclerosis (cor = -0.19, SE = 0.07, FDR < 0.06). These results highlight the shared genetic architecture underlying central mechanism in AD and other neuroinflammatory disorders [83, 84]. Recent GWAS and pathway analyses have emphasized the crucial role of the innate immune system and neuroinflammation in the pathogenesis of AD [85, 86]. To that end, targeting neuroinflammation by modulating different phagocytic receptors e.g. CD33 inhibition [87] and/or TREM2 activation [88, 89] have been suggested as valuable therapeutic strategies to enhance neuroprotective Page 15 of 20

microglia and reduce neuroinflammation, which is crucial for preventing and treating AD [85].

Similar to other GWAS, the present study also bears some important limitations. Although we have compiled and leveraged the largest collection of amyloid PET datasets reported-to-date, the sample size for all populations were not comparable. Significantly reduced sample size from the AFR and ASN ancestries might be the most profound hurdle in achieving genome-wide significance for loci with small effect sizes across multi-ethnic meta-analysis. Nevertheless, we showed that some of our suggestive and nominally significant loci have the same direction of allelic effects for established AD risk associated variants, suggesting that we might have achieved genome-wide significance with a relatively larger sample size. One potential reason for relatively lower sample size is the difficulty in obtaining the amyloid PET data through scanning due to limited availability of scanning platform in the under-developed countries and their lesser representation in developed countries where such facilities are readily available. Another significant constraint was the cis-eQTL dataset that was obtained from GTEx database which only represents cognitively normal individuals and not those diagnosed with AD. As more amyloid PET imaging data are obtained by different centers with larger population sizes, future studies leveraging these larger samples sizes will have better potential for validating existing findings and identifying additional genes associated with brain amyloidosis.

## Conclusion

We have performed the largest reported to-date amyloid PET GWAS (N=13,409) that has confirmed the previously known association of the *APOE* locus with brain amyloidosis. In addition to recapitulating the established associations, we have identified novel variants in the *ABCA7*, *CR1*, and *FERMT2* regions as well as sexspecific variants that affect amyloid deposition. We have employed a combination of genetic and functional analytic approaches for identifying putative candidate genes that warrant follow-up genetic and functional studies to confirm their role in brain amyloidosis. This study highlights the importance and need of large-scale genetic studies focusing on brain amyloidosis in diverse populations for finding universal candidate therapeutic targets for AD.

#### Abbreviations

AD

Aβ GW/

PiB

MAR

Knic

	Alzheimer's disease
	Amyloid beta
AS	Genome-wide association studies
	Pittsburgh compound-B
P	Memory and aging project
ght-ADRC	Knight alzheimer's disease research center

ADNI DIAN A4	Alzheimer's disease neuroimaging initiative Dominantly inherited alzheimer network Anti-amyloid treatment in asymptomatic alzheimer's
/ ( )	disease
ADNI-DOD	adni department of defense
AIBL	Australian imaging, biomarkers and lifestyle
HABS	Harvard aging brain study
UPitt	University of pittsburgh
MCSA	Mayo clinic study of aging
WRAP	Wisconsin registry for alzheimer's prevention
BACS	Berkeley aging cohort study
CDR	Clinical dementia rating
PCs	Principal components
SUVR	Standardize uptake value ratios
SD	Standard deviation
MAF	Minor allele frequency
QC	Quality control
IBD	Identity-by-descent
PCA	Principal component analysis
NHW	Non-Hispanic Whites
AFR	American Africans
ASN	Asians
StdErr	Standard error
GNOVA	Genetic covariance analyzer
GTEx	Genotype-tissue expression
PRS	Polygenic risk score
LD	Linkage disequilibrium
QTL	Quantitative trait loci
sQTLs	Single-tissue splicing QTLs
APOE	Apolipoprotein E
CAA	Cerebral amyloid angiopathy
CO	Controls
CR1	Complement receptor 1
HDL	High-density lipoprotein
TOMM40	Translocase of outer mitochondrial membrane 40
	homolog
EHF	ETS homologous factor
ABCA7	ATP-binding cassette sub-family a member 7
BIN1	Bridging integrator 1
ANK3	Ankyrin 3
TREM2	Triggering receptor expressed on myeloid cells 2
CLU	Clusterin Arrestantia latara lateralia
ALS	Amyotrophic lateral sclerosis
MCI	Mild cognitive impairment

## **Supplementary Information**

The online version contains supplementary material available at https://doi. org/10.1186/s40478-023-01563-4.

Additional file 1. Supplementary tables and figures

#### Acknowledgements

We thank all the participants and their families, as well as the involved cohorts, institutions, and their staff. The recruitment and clinical characterization of research participants at Washington University were supported by NIH P30 AG066444 (JCM), P01 AG03991 (JCM), and P01 AG026276 (JCM). This work was supported by access to equipment made possible by the Hope Center for Neurological Disorders, the Neurogenomics and Informatics Center (NGI: https://neurogenomics.wustl.edu/) and the Departments of Neurology and Psychiatry at Washington University School of Medicine.

**DIAN resources**: Data collection and sharing for this project was supported by The Dominantly Inherited Alzheimer Network (DIAN, U19 AG032438) funded by the National Institute on Aging (NIA), the Alzheimer's Association (SG-20-690363-DIAN), the German Center for Neurodegenerative Diseases (DZNE), Raul Carrea Institute for Neurological Research (FLENI), Partial support by the Research and Development Grants for Dementia from Japan Agency for Medical Research and Development, AMED, and the Korea Health Technology R&D Project through the Korea Health Industry Development Institute (KHIDI), Spanish Institute of Health Carlos III (ISCIII), Canadian Institutes of Health Research (CIHR), Canadian Consortium of Neurodegeneration and Aging, Brain Canada Foundation, and Fonds de Recherche du Québec – Santé. This manuscript has been reviewed by DIAN Study investigators for scientific content and consistency of data interpretation with previous DIAN Study publications. We acknowledge the altruism of the participants and their families and contributions of the DIAN research and support staff at each of the participating sites for their contributions to this study.

#### ADNI acknowledgement

Data collection and sharing for this project was funded by the Alzheimer's Disease Neuroimaging Initiative (ADNI) (National Institutes of Health Grant U01 AG024904) and DOD ADNI (Department of Defense award number W81XWH-12-2-0012). ADNI is funded by the National Institute on Aging, the National Institute of Biomedical Imaging and Bioengineering, and through generous contributions from the following: AbbVie, Alzheimer's Association; Alzheimer's Drug Discovery Foundation; Araclon Biotech; BioClinica, Inc.; Biogen; Bristol-Myers Squibb Company; CereSpir, Inc.; Cogstate; Eisai Inc.; Elan Pharmaceuticals, Inc.; Eli Lilly and Company; EuroImmun; F. Hoffmann-La Roche Ltd and its affiliated company Genentech, Inc.; Fujirebio; GE Healthcare; IXICO Ltd.; Janssen Alzheimer Immunotherapy Research & Development, LLC.; Johnson & Johnson Pharmaceutical Research & Development LLC.; Lumosity; Lundbeck; Merck & Co., Inc.; Meso Scale Diagnostics, LLC.; NeuroRx Research; Neurotrack Technologies; Novartis Pharmaceuticals Corporation; Pfizer Inc.; Piramal Imaging; Servier; Takeda Pharmaceutical Company; and Transition Therapeutics. The Canadian Institutes of Health Research is providing funds to support ADNI clinical sites in Canada. Private sector contributions are facilitated by the Foundation for the National Institutes of Health (www.fnih. org). The grantee organization is the Northern California Institute for Research and Education, and the study is coordinated by the Alzheimer's Therapeutic Research Institute at the University of Southern California. ADNI data are disseminated by the Laboratory for Neuro Imaging at the University of Southern California.

#### MCSA acknowledgement

Mayo Clinic Study of Aging data was acquired with the support of NIH grants U01 AG006786, R01 AG56366, P50 AG016574, P30 AG062677, R37 AG011378, R01 AG041851, R01 AG034676; GHR Foundation, and the Mayo Foundation for Medical Education and Research.

### **GEMS** acknowledgement

This study was supported by grant U01 AT000162 from the National Center for Complementary and Alternative Medicine and the Office of Dietary Supplements, and grant P01 AG025204 from the National Institute on Aging. We would also like to acknowledge the scientific contributions of GEMS investigators from the Pittsburgh site; Lewis H Kuller, William Klunk, Chester Mathis and Howard Aizenstein.

#### A4 acknowledgement

The A4 Study is a secondary prevention trial in preclinical Alzheimer's disease, aiming to slow cognitive decline associated with brain amyloid accumulation in clinically normal older individuals. The A4 Study is funded by a public-private-philanthropic partnership, including funding from the National Institutes of Health-National Institute on Aging (R01 AG063689, U24 AG057437) Eli Lilly and Company, Alzheimer's Association, Accelerating Medicines Partnership, GHR Foundation, an anonymous foundation and additional private donors, with in-kind support from Avid and Cogstate. The companion observational Longitudinal Evaluation of Amyloid Risk and Neurodegeneration (LEARN) Study is funded by the Alzheimer's Association and GHR Foundation. The A4 and LEARN Studies are led by Dr. Reisa Sperling at Brigham and Women's Hospital, Harvard Medical School and Dr. Paul Aisen at the Alzheimer's Therapeutic Research Institute (ATRI), University of Southern California. The A4 and LEARN Studies are coordinated by ATRI at the University of Southern California, and the data are made available through the Laboratory for Neuro Imaging at the University of Southern California. The participants screening for the A4 Study provided permission to share their de-identified data in order to advance the quest to find a successful treatment for Alzheimer's disease. We would like to acknowledge the dedication of all the participants, the site personnel, and all of the partnership team members who continue to make the A4 and LEARN Studies possible. The complete A4 Study Team list is available on: a4study.org/ a4-study-team.

Data used in preparation of this article were obtained from the Alzheimer's Disease Neuroimaging Initiative (ADNI) database (adni.loni.usc.edu). As such, the investigators within the ADNI contributed to the design and implementation of ADNI and/or provided data but did not participate in analysis or writing of this report. A complete listing of ADNI investigators can be found at: http://adni.loni.usc.edu/wp-content/uploads/how\_to\_apply/ADNI\_Acknowledg ement\_List.pdf

The amyloid PET and data were acquired during screening for the A4 Study and LEARN by the A4 Study Team. Full team authorship list available at https://a4study.org/

**AIBL acknowledgement** Data used in the preparation of this article was obtained from the Australian Imaging Biomarkers and Lifestyle flagship study of ageing (AIBL) funded by the Commonwealth Scientific and Industrial Research Organisation (CSIRO). The AIBL researchers, unless listed, contributed data but did not participate in the analysis or writing of this report. AIBL researchers are listed at www.aibl.csiro.au.

#### Author contributions

MA Conceptualization, Formal analysis, Writing – original draft, Writing – review & editing. CC: Conceptualization, Funding acquisition, Supervision, Validation, Writing – review & editing. DBA and LD: Data curation, Formal analysis, Review & editing. PG, DW, JT, MVF, TCW, CLS, QY, ASB, RQ, GC, BG, TLSB, CX, RJB, CMK, EM, AG, SS, RPM, RAS, RFB, KAJ, HW, SJ, HK, SWS, HJK, EM, SML, KF, MIK, PV, VKR, HY, WJJ, AW, HSRR, AM, CD, SD, OL, STD, FT, MWN, TJH, YJS: Data curation, Review & editing. JCM: Funding acquisition, Project administration, Review & editing.

#### Funding

This work was supported by grants from the National Institutes of Health, R01 AG044546 (CC), P01 AG003991 (CC, JCM), RF1 AG053303 (CC), RF1 AG058501 (CC), U01 AG058922 (CC), R01 AG064877 (MIK), P30 AG066468 (MIK), U01 AT000162 (MIK), K23 AG062750 (HY), and the Chan Zuckerberg Initiative (CZI), the Michael J. Fox Foundation (LI, CC), the Department of Defense (LI-W81XWH2010849), and the Alzheimer's Association Zenith Fellows Award (ZEN-22-848604, awarded to CC). Dr. Mathis and Dr. Klunk receive royalty payments from GE Healthcare (indirect through a license agreement with the University of Pittsburgh). GE Healthcare holds a license agreement with the University of Pittsburgh based on the technology described in this manuscript. Dr. Mathis and Dr Klunk are a co-inventor of PiB and, as such, has a financial interest in this license agreement. GE Healthcare provided no grant support for this study and had no role in the design or interpretation of results or preparation of this manuscript. All other authors have no conflicts of interest with this work. CC has received research support from: GSK and EISAI. The funders of the study had no role in the collection, analysis, or interpretation of data; in the writing of the report; or in the decision to submit the paper for publication. CC is a member of the advisory board of Vivid Genomics and Circular Genomics and owns Stocks. Neither Dr. Morris nor his family owns stock or has equity interest (outside of mutual funds or other externally directed accounts) in any pharmaceutical or biotechnology company.

#### Availability of data and materials

All data generated in this study are included in this published article.

## Declarations

#### Ethics approval and consent to participate

Ethics approval for every individual cohort was obtained from the respective Institutional Review Boards and research was carried out in accordance with the approved protocols (WUSTL IRB approval 201109148). Written informed consent was obtained from participants or their family members and all participating institutions approved the study design.

#### Consent for publication

All authors have approved the contents of this manuscript and provided consent for publication.

#### **Competing interests**

The authors declare that they no competing interests.

#### Author details

Department of Psychiatry, Washington University, St. Louis, MO 63110, USA. <sup>2</sup>NeuroGenomics and Informatics, Washington University, St. Louis, MO 63110, USA. <sup>3</sup>Vanderbilt Memory and Alzheimer's Center, Vanderbilt University School of Medicine, Nashville, TN, USA. <sup>4</sup>Glenn Biggs Institute for Alzheimer's and Neurodegenerative Diseases, UT Health, San Antonio, TX 78229, USA. <sup>5</sup>Department of Biostatistics, Boston University School of Public Health, Boston, MA, USA. <sup>6</sup>Department of Neurology, Boston University School of Medicine, Boston, MA, USA. <sup>7</sup>Boston University, Boston, MA, USA. <sup>8</sup>Knight Alzheimer's Disease Research Center, Washington University, St Louis, MO, USA. <sup>9</sup>Mallinckrodt Institute of Radiology, Washington University, St Louis, MO, USA. <sup>10</sup>Department of Neurology, Washington University, St Louis, MO, USA. <sup>11</sup>Department of Neuroscience, Ronald M. Loeb Center for Alzheimer's Disease, Icahn School of Medicine at Mount Sinai, New York, NY, USA. <sup>12</sup>Framingham Heart Study, Framingham, MA, USA. <sup>13</sup>Boston University School of Medicine, Boston, MA, USA. <sup>14</sup>The Department of Neurology, Columbia University, New York, NY, USA. <sup>15</sup>Department of Neurology, Harvard Medical School, Boston, MA, USA. <sup>16</sup>Brigham and Women's Hospital and Department of Neurology, Massachusetts General Hospital, Harvard Medical School, Boston, MA, USA. <sup>17</sup>Athinoula A. Martinos Center for Biomedical Imaging, Charlestown, MA, USA. <sup>18</sup>Massa chusetts General Hospital, Harvard Medical School, Boston, MA, USA. <sup>19</sup>Department of Digital Health, Samsung Medical Center, SAIHST, Sungkyunkwan University, Seoul, Republic of Korea.<sup>20</sup>Department of Neurology, Dongguk University Ilsan Hospital, Dongguk University College of Medicine, Goyang, Republic of Korea.<sup>21</sup>Department of Neurology, Samsung Medical Center, Sungkyunkwan University School of Medicine, Seoul, Republic of Korea. <sup>22</sup>Department of Neurology and Neurological Sciences, Stanford University, Stanford, CA, USA. <sup>23</sup>Centre for Precision Health, Edith Cowan University, 270 Joondalup Dr, Joondalup, WA 6027, Australia.<sup>24</sup>Department of Human Genetics, University of Pittsburgh, Pittsburgh, PA, USA.<sup>25</sup>Department of Radiology, Mayo Clinic-Minnesota, Rochester, MN 55905, USA. <sup>26</sup>Department of Neurology, Mayo Clinic-Minnesota, Rochester, MN 55905, USA. <sup>27</sup>Department of Neurology, Brigham and Women's Hospital, Boston, MA, USA. <sup>28</sup>Department of Neurology, Massachusetts General Hospital, Boston, MA, USA.<sup>29</sup>Harvard Medical School, Boston, MA, USA. <sup>30</sup>Broad Institute of Harvard and MIT, Cambridge, USA. <sup>31</sup>Wisconsin Alzheimer's Institute, Madison, WI, USA. <sup>32</sup>UMR 1219, University of Bordeaux, INSERM, Bordeaux Population Health Research Centre, Team ELEANOR, 33000 Bordeaux, France. <sup>33</sup>Department of Neurology, Boston University School of Medicine, Boston, MA 2115, USA. <sup>34</sup>Department of Neurology, CHU de Bordeaux, 33000 Bordeaux, France. <sup>35</sup>Department of Neurology, University of Pittsburgh, Pittsburgh, PA, USA. <sup>36</sup>Department of Neurology and McKnight Brain Institute, University of Florida, Gainesville, FL, USA. <sup>37</sup>Neurogenomics, Genetics-Guided Dementia Discovery, Eisai, Inc, Cambridge, MA, USA. <sup>38</sup>Hope Center for Neurologic Diseases, Washington University, St. Louis, MO 63110, USA. <sup>39</sup>Department of Genetics, Washington University School of Medicine, St Louis, MO 63110, USA.

## Received: 27 January 2023 Accepted: 5 April 2023 Published online: 26 April 2023

#### References

- Gatz M, Reynolds CA, Fratiglioni L, Johansson B, Mortimer JA, Berg S et al (2006) Role of genes and environments for explaining Alzheimer disease. Arch Gen Psych United States 63:168–174
- Jansen IE, Savage JE, Watanabe K, Bryois J, Williams DM, Steinberg S et al (2019) Genome-wide meta-analysis identifies new loci and functional pathways influencing Alzheimer's disease risk. Nat Genet [Internet]. 51:404–413. https://doi.org/10.1038/s41588-018-0311-9
- 3. Kunkle BW, Grenier-Boley B, Sims R, Bis JC, Damotte V, Naj AC et al (2019) Genetic meta-analysis of diagnosed Alzheimer's disease identifies new risk loci and implicates A $\beta$ , tau, immunity and lipid processing. Nat Genet [Internet]. 51:414–430. https://doi.org/10.1038/s41588-019-0358-2
- Schwartzentruber J, Cooper S, Liu JZ, Barrio-Hernandez I, Bello E, Kumasaka N et al (2021) Genome-wide meta-analysis, fine-mapping and integrative prioritization implicate new Alzheimer's disease risk genes. Nat Genet 53:392–402
- 5. Bellenguez C, Küçükali F, Jansen IE, Kleineidam L, Moreno-Grau S, Amin N et al (2022) New insights into the genetic etiology of Alzheimer's disease

and related dementias. Nat Genet [Internet]. 54:412–436. https://doi.org/ 10.1038/s41588-022-01024-z

- Ridge PG, Hoyt KB, Boehme K, Mukherjee S, Crane PK, Haines JL et al (2016) Assessment of the genetic variance of late-onset Alzheimer's disease. Neurobiol Aging 41:200.e13-200.e20
- Hyman BT, Phelps CH, Beach TG, Bigio EH, Cairns NJ, Carrillo MC et al (2012) National Institute on Aging-Alzheimer's Association guidelines for the neuropathologic assessment of Alzheimer's disease. Alzheimers Dement 8:1–13
- Jack CRJ, Therneau TM, Wiste HJ, Weigand SD, Knopman DS, Lowe VJ et al (2016) Transition rates between amyloid and neurodegeneration biomarker states and to dementia: a population-based, longitudinal cohort study. Lancet Neurol 15:56–64
- Burnham SC, Bourgeat P, Doré V, Savage G, Brown B, Laws S et al (2016) Clinical and cognitive trajectories in cognitively healthy elderly individuals with suspected non-Alzheimer's disease pathophysiology (SNAP) or Alzheimer's disease pathology: a longitudinal study. Lancet Neurol England 15:1044–1053
- 10. Sturchio A, Dwivedi AK, Young CB, Malm T, Marsili L, Sharma JS et al (2021) High cerebrospinal amyloid- $\beta$  42 is associated with normal cognition in individuals with brain amyloidosis. EClinicalMedicine 38:100988
- Yan Q, Nho K, Del-Aguila J, Wang X, Risacher S, Fan K, et al. (2021) Genome-wide association study of brain amyloid deposition as measured by Pittsburgh Compound-B (PiB)-PET imaging. Mol Psychiatry [Internet]. Mol Psychiatry, 26:309–321. Available from: https://pubmed. ncbi.nlm.nih.gov/30361487/
- Camus V, Payoux P, Barré L, Desgranges B, Voisin T, Tauber C et al (2012) Using PET with 18F-AV-45 (florbetapir) to quantify brain amyloid load in a clinical environment. Eur J Nucl Med Mol Imaging 39:621–631
- Vacher M, Porter T, Villemagne VL, Milicic L, Peretti M, Fowler C et al (2019) Validation of a priori candidate Alzheimer's disease SNPs with brain amyloid-beta deposition. Sci Rep [Internet]. 9:17069. https://doi.org/10. 1038/s41598-019-53604-5
- Li J, Zhang Q, Chen F, Yan J, Kim S, Wang L et al (2015) Genetic interactions explain variance in cingulate amyloid burden: an AV-45 PET genome-wide association and interaction study in the ADNI cohort. Biomed Res Int 2015;647389
- Raghavan NS, Dumitrescu L, Mormino E, Mahoney ER, Lee AJ, Gao Y et al (2020) Association between common variants in RBFOX1, an RNAbinding protein, and brain amyloidosis in early and preclinical alzheimer disease. JAMA Neurol [Internet]. 77:1288–1298. https://doi.org/10.1001/ jamaneurol.2020.1760
- Deming Y, Li Z, Kapoor M, Harari O, Del-Aguila JL, Black K et al (2017) Genome-wide association study identifies four novel loci associated with Alzheimer's endophenotypes and disease modifiers. Acta Neuropathol. https://doi.org/10.1007/s00401-017-1685-y
- 17. Morris JC, Roe CM, Xiong C, Fagan AM, Goate AM, Holtzman DM et al (2010) APOE predicts amyloid-beta but not tau Alzheimer pathology in cognitively normal aging. Ann Neurol 67:122–131
- Swaminathan S, Shen L, Risacher SL, Yoder KK, West JD, Kim S et al (2012) Amyloid pathway-based candidate gene analysis of [(11)C]PiB-PET in the Alzheimer's Disease Neuroimaging Initiative (ADNI) cohort. Brain Imaging Behav 6:1–15
- Prescott JW, Doraiswamy PM, Gamberger D, Benzinger T, Petrella JR (2021) Diffusion Tensor MRI Structural Connectivity and PET Amyloid Burden in Preclinical Autosomal Dominant Alzheimer Disease: The DIAN Cohort. Radiology 302:143–150. https://doi.org/10.1148/radiol.20212 10383
- Sperling RA, Rentz DM, Johnson KA, Karlawish J, Donohue M, Salmon DP et al (2014) The A4 study: stopping AD before symptoms begin? Sci Transl Med. 6:228fs13
- Rowe CC, Ellis KA, Rimajova M, Bourgeat P, Pike KE, Jones G et al (2010) Amyloid imaging results from the Australian Imaging, Biomarkers and Lifestyle (AIBL) study of aging. Neurobiol Aging United States 31:1275–1283
- Dagley A, LaPoint M, Huijbers W, Hedden T, McLaren DG, Chatwal JP et al (2017) Harvard aging brain study: dataset and accessibility. Neuroimage 144:255–258
- 23. Mathis CA, Kuller LH, Klunk WE, Snitz BE, Price JC, Weissfeld LA et al (2013) In vivo assessment of amyloid- $\beta$  deposition in nondemented very elderly subjects. Ann Neurol 73:751–761

- 24. Gallet Q, Bouteloup V, Locatelli M, Habert M-O, Chupin M, Delrieu J et al (2022) Benzodiazepine use and neuroimaging markers of Alzheimer's disease in nondemented older individuals: an MRI and 18F Florbetapir PET study in the MEMENTO cohort. Neuropsychopharmacology 47:1114–1420. https://doi.org/10.1038/s41386-021-01246-5
- 25. Ramanan VK, Wang X, Przybelski SA, Raghavan S, Heckman MG, Batzler A et al (2020) Variants in PPP2R2B and IGF2BP3 are associated with higher tau deposition. Brain Commun. 2:fcc159. https://doi.org/10.1093/brain comms/fcaa159
- Johnson SC, Koscik RL, Jonaitis EM, Clark LR, Mueller KD, Berman SE et al (2018) The wisconsin registry for alzheimer's prevention: a review of findings and current directions. Alzheimer's Dement (Amsterdam, Netherlands) 10:130–142
- Ossenkoppele R, Madison C, Oh H, Wirth M, van Berckel BNM, Jagust WJ (2014) Is verbal episodic memory in elderly with amyloid deposits preserved through altered neuronal function? Cereb Cortex 24:2210–2218
- Kim H-R, Jung S-H, Kim J, Jang H, Kang SH, Hwangbo S et al (2021) Identifying novel genetic variants for brain amyloid deposition: a genome-wide association study in the Korean population. Alzheimers Res Ther. 13:117. https://doi.org/10.1186/s13195-021-00854-z
- 29. Gonzales MM, Samra J, O'Donnell A, Mackin RS, Salinas J, Jacob ME et al (2021) Association of midlife depressive symptoms with regional amyloid- $\beta$  and tau in the framingham heart study. J Alzheimers Dis 82:249–260
- Loh P-R, Danecek P, Palamara PF, Fuchsberger C, A Reshef Y, K Finucane H et al (2016) Reference-based phasing using the Haplotype Reference Consortium panel. Nat Genet 48:1443–1448. https://doi.org/10.1038/ng. 3679
- Howie B, Fuchsberger C, Stephens M, Marchini J, Abecasis GR (2012) Fast and accurate genotype imputation in genome-wide association studies through pre-phasing. Nat Genet. 44(8):955–959
- Chang CC, Chow CC, Tellier LCAM, Vattikuti S, Purcell SM, Lee JJ (2015) Second-generation PLINK: Rising to the challenge of larger and richer datasets. Gigascience. https://doi.org/10.1186/s13742-015-0047-8
- Willer CJ, Li Y, Abecasis GR (2010) METAL: fast and efficient meta-analysis of genomewide association scans. Bioinformatics 26:2190–2191
- R Core Team (2021) R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna, Austria. https:// www.R-project.org/
- Skol AD, Scott LJ, Abecasis GR, Boehnke M (2006) Joint analysis is more efficient than replication-based analysis for two-stage genome-wide association studies. Nat Genet. 38:209–213. https://doi.org/10.1038/ ng1706
- Ma C, Blackwell T, Boehnke M, Scott LJ (2013) Recommended joint and meta-analysis strategies for case-control association testing of single lowcount variants. Genet Epidemiol 37:539–550
- Teissier M, Sanchez MP, Boussaha M, Barbat A, Hoze C, Robert-Granie C, et al. (2018) Use of meta-analyses and joint analyses to select variants in whole genome sequences for genomic evaluation: An application in milk production of French dairy cattle breeds. J Dairy Sci [Internet]. 101:3126–3139. https://www.sciencedirect.com/science/article/pii/S0022 030218300857
- Lu Q, Li B, Ou D, Erlendsdottir M, Powles RL, Jiang T et al (2017) A powerful approach to estimating annotation-stratified genetic covariance via GWAS summary statistics. Am J Hum Genet 101:939–964
- 39. Wallace C (2021) A more accurate method for colocalisation analysis allowing for multiple causal variants. PLoS Genet 17:e1009440
- Brouwers N, Van Cauwenberghe C, Engelborghs S, Lambert J-C, Bettens K, Le Bastard N et al (2012) Alzheimer risk associated with a copy number variation in the complement receptor 1 increasing C3b/C4b binding sites. Mol Psychiatry 17:223–233. https://doi.org/10.1038/mp.2011.24
- Euesden J, Lewis CM, O'Reilly PF (2015) PRSice: polygenic risk score software. Bioinformatics 31:1466–1468
- 42. Lonsdale J, Thomas J, Salvatore M, Phillips R, Lo E, Shad S et al (2013) The genotype-tissue expression (GTEx) project. Nat Genet. 45:580–585. https://doi.org/10.1038/ng.2653
- Fonseca MI, Chu S, Pierce AL, Brubaker WD, Hauhart RE, Mastroeni D et al (2016) Analysis of the putative role of CR1 in alzheimer's disease: genetic association, expression and Function. PLoS One 11:1–21. https://doi.org/ 10.1371/journal.pone.0149792

- Yan Y, Zhao A, Qui Y, Li Y, Yan R, Wang Y, et al. (2020) Genetic Association of FERMT2, HLA-DRB1, CD2AP, and PTK2B Polymorphisms With Alzheimer's Disease Risk in the Southern Chinese Population. Front Aging Neurosci. 12. Available from: https://www.frontiersin.org/articles/https://doi.org/10. 3389/fnagi.2020.00016
- Zhang B, Gaiteri C, Bodea L-G, Wang Z, McElwee J, Podtelezhnikov AA, et al. (2013) Integrated systems approach identifies genetic nodes and networks in late-onset Alzheimer's disease. Cell, 153:707–720 https:// pubmed.ncbi.nlm.nih.gov/23622250
- 46. Sullivan SE, Liao M, Smith RV, White C, Lagomarsino VN, Xu J et al (2019) Candidate-based screening via gene modulation in human neurons and astrocytes implicates FERMT2 in Aβ and TAU proteostasis. Hum Mol Genet England 28:718–735
- Greenwood AK, Gockley J, Daily K, Aluthgamage D, Leanza Z, Sieberts SK et al (2020) Agora: An open platform for exploration of Alzheimer's disease evidence. Alzheimer's Dement 16:e046129. https://doi.org/10. 1002/alz.046129
- Sakae N, Liu C-C, Shinohara M, Frisch-Daiello J, Ma L, Yamazaki Y et al (2016) ABCA7 deficiency accelerates amyloid-β generation and alzheimer's neuronal pathology. J Neurosci Off J Soc Neurosci 36:3848–3859
- Logge W, Cheng D, Chesworth R, Bhatia S, Garner B, Kim WS et al (2012) Role of Abca7 in mouse behaviours relevant to neurodegenerative diseases. PLoS ONE 7:e45959
- De Roeck A, Van Broeckhoven C, Sleegers K (2019) The role of ABCA7 in Alzheimer's disease: evidence from genomics, transcriptomics and methylomics. Acta Neuropathol 138:201–220
- Farfel JM, Yu L, Buchman AS, Schneider JA, De Jager PL, Bennett DA (2016) Relation of genomic variants for Alzheimer disease dementia to common neuropathologies. Neurology 87:489–496
- Lamar M, Yu L, Rubin LH, James BD, Barnes LL, Farfel JM et al (2019) APOE genotypes as a risk factor for age-dependent accumulation of cerebrovascular disease in older adults. Alzheimer's Dement 15:258–266. https:// doi.org/10.1016/j.jalz.2018.08.007
- Li Z, Shue F, Zhao N, Shinohara M, Bu G (2020) APOE2: protective mechanism and therapeutic implications for Alzheimer's disease. Mol Neurodegener 15:63
- Serrano-Pozo A, Qian J, Monsell SE, Betensky RA, Hyman BT (2015) APOEc2 is associated with milder clinical and pathological Alzheimer disease. Ann Neurol 77:917–929
- Yang J, Ferreira T, Morris AP, Medland SE, Madden PAF, Heath AC et al (2012) Conditional and joint multiple-SNP analysis of GWAS summary statistics identifies additional variants influencing complex traits. Nat Genet 44:369–75. https://doi.org/10.1038/ng.2213
- McLaren W, Gil L, Hunt SE, Riat HS, Ritchie GRS, Thormann A et al (2016) The Ensembl Variant Effect Predictor. Genome Biol 17:122. https://doi. org/10.1186/s13059-016-0974-4
- 57. Association A (2019) Alzheimer's disease facts and figures. Alzheimer's Dement, 15:321–387, https://doi.org/10.1016/j.jalz.2019.01.010
- Dumitrescu L, Mahoney ER, Mukherjee S, Lee ML, Bush WS, Engelman CD et al (2020) Genetic variants and functional pathways associated with resilience to Alzheimer's disease. Brain 143:2561–2575. https://doi.org/10. 1093/brain/awaa209
- Shen L, Kim S, Risacher SL, Nho K, Swaminathan S, West JD et al (2010) Whole genome association study of brain-wide imaging phenotypes for identifying quantitative trait loci in MCI and AD: A study of the ADNI cohort. Neuroimage 53:1051–1063
- Ferencz B, Laukka EJ, Lövdén M, Kalpouzos G, Keller L, Graff C et al (2013) The influence of APOE and TOMM40 polymorphisms on hippocampal volume and episodic memory in old age. Front Hum Neurosci 7:198
- Siddarth P, Burggren AC, Merrill DA, Ercoli LM, Mahmood Z, Barrio JR et al (2018) Longer TOMM40 poly-T variants associated with higher FDDNP-PET medial temporal tau and amyloid binding. PLoS ONE 13:e0208358
- Cruchaga C, Nowotny P, Kauwe JSK, Ridge PG, Mayo K, Bertelsen S et al (2011) Association and expression analyses with single-nucleotide polymorphisms in TOMM40 in Alzheimer disease. Arch Neurol United States 68:1013–1019
- 63. Oatman SR, Reddy JS, Quicksall Z, Carrasquillo MM, Wang X, Liu C-C et al (2023) Genome-wide association study of brain biochemical phenotypes reveals distinct genetic architecture of Alzheimer's disease related proteins. Mol Neurodegener. 18:2. https://doi.org/10.1186/ s13024-022-00592-2

- 64. Seshadri S, Fitzpatrick AL, Ikram MA, DeStefano AL, Gudnason V, Boada M et al (2010) Genome-wide analysis of genetic loci associated with Alzheimer disease. JAMA United States 303:1832–1840
- 65. Cruchaga C, Kauwe JSK, Harari O, Jin SC, Cai Y, Karch CM et al (2013) GWAS of cerebrospinal fluid tau levels identifies risk variants for Alzheimer's disease. Neuron United States 78:256–268
- 66. Mafimoghaddam S, Wang L, Gorijala P, Timsina J, Nykanen N, Wang F et al (2022) GWAS for CSF TREM2 levels identify new variants implicated on TREM2 biology and Alzheimer disease. Alzheimer's Dement 18:e062517. https://doi.org/10.1002/alz.062517
- Arboleda-Velasquez JF, Lopera F, O'Hare M, Delgado-Tirado S, Marino C, Chmielewska N et al (2019) Resistance to autosomal dominant Alzheimer's disease in an APOE3 Christchurch homozygote: a case report. Nat Med. 25:1680–1683. https://doi.org/10.1038/s41591-019-0611-3
- Nho K, Kim S, Horgusluoglu E, Risacher SL, Shen L, Kim D et al (2017) Association analysis of rare variants near the APOE region with CSF and neuroimaging biomarkers of Alzheimer's disease. BMC Med Genomics 10:29
- Farrer LA, Cupples LA, Haines JL, Hyman B, Kukull WA, Mayeux R et al (1997) Effects of age, sex, and ethnicity on the association between apolipoprotein E genotype and Alzheimer disease. A meta-analysis. APOE and Alzheimer Disease Meta Analysis Consortium. JAMA 278(16):1349–1356. https://doi.org/10.1001/jama.1997.0355016006 9041
- van Dyck CH, Swanson CJ, Aisen P, Bateman RJ, Chen C, Gee M et al (2022) Lecanemab in Early Alzheimer's Disease. N Engl J Med. https:// doi.org/10.1056/NEJMoa2212948
- Steinberg S, Stefansson H, Jonsson T, Johannsdottir H, Ingason A, Helgason H et al (2015) Loss-of-function variants in ABCA7 confer risk of Alzheimer's disease. Nat Genet. 47:445–457. https://doi.org/10.1038/ ng.3246
- Vasquez JB, Fardo DW, Estus S (2013) ABCA7 expression is associated with Alzheimer's disease polymorphism and disease status. Neurosci Lett 556:58–62
- Chapuis J, Hansmannel F, Gistelinck M, Mounier A, Van Cauwenberghe C, Kolen KV et al (2013) Increased expression of BIN1 mediates Alzheimer genetic risk by modulating tau pathology. Mol Psychiatry 18:1225–1234
- Jonsson T, Stefansson H, Steinberg S, Jonsdottir I, Jonsson PV, Snaedal J et al (2013) Variant of TREM2 associated with the risk of Alzheimer's disease. N Engl J Med 368:107–116
- Guerreiro R, Wojtas A, Bras J, Carrasquillo M, Rogaeva E, Majounie E, et al. (2012) TREM2 variants in Alzheimer's disease. N Engl J Med, 368:117–127 https://pubmed.ncbi.nlm.nih.gov/23150934
- Foster EM, Dangla-Valls A, Lovestone S, Ribe EM, Buckley NJ (2019) Clusterin in Alzheimer's Disease: mechanisms, genetics, and lessons from other pathologies. Front Neurosci 13:164
- Lunnon K, Smith R, Hannon E, De Jager PL, Srivastava G, Volta M et al (2014) Methylomic profiling implicates cortical deregulation of ANK1 in Alzheimer's disease. Nat Neurosci 17:1164–1170
- Hayashi M, Abe-Dohmae S, Okazaki M, Ueda K, Yokoyama S (2005) Heterogeneity of high density lipoprotein generated by ABCA1 and ABCA7. J Lipid Res United States 46:1703–1711
- 79. Feringa FM, van der Kant R (2021) Cholesterol and Alzheimer's Disease; from risk genes to pathological effects. Front Aging Neurosci 13:690372
- Gu M-L, Zhou X-X, Ren M-T, Shi K-D, Yu M-S, Jiao W-R et al (2020) Blockage of ETS homologous factor inhibits the proliferation and invasion of gastric cancer cells through the c-Met pathway. World J Gastroenterol United States 26:7497–7512
- Area-Gomez E, Del Carmen Lara Castillo M, Tambini MD, Guardia-Laguarta C, de Groof AJC, Madra M et al (2012) Upregulated function of mitochondria-associated ER membranes in Alzheimer disease. EMBO J 31:4106–4123
- Smith CD, Van Eldik LJ, Jicha GA, Schmitt FA, Nelson PT, Abner EL et al (2020) Brain structure changes over time in normal and mildly impaired aged persons. AIMS Neurosci 7:120–135
- Akiyama H, Barger S, Barnum S, Bradt B, Bauer J, Cole GM et al (2000) Inflammation and Alzheimer's disease. Neurobiol Aging United States 21:383–421

- Kinney JW, Bemiller SM, Murtishaw AS, Leisgang AM, Salazar AM, Lamb BT (2018) Inflammation as a central mechanism in Alzheimer's disease. Alzheimer's Dement 4:575–590
- Griciuc A, Tanzi RE (2021) The role of innate immune genes in Alzheimer's disease. Curr Opin Neurol [Internet]. 34. Available from: https://journ als.lww.com/co-neurology/Fulltext/2021/04000/The\_role\_of\_innate\_ immune\_genes\_in\_Alzheimer\_s.13.aspx
- Griciuc A, Patel S, Federico AN, Choi SH, Innes BJ, Oram MK et al (2019) TREM2 acts downstream of CD33 in modulating microglial pathology in alzheimer's disease. Neuron United States 103:820-835.e7
- Zhao L (2019) CD33 in Alzheimer's Disease biology, pathogenesis, and therapeutics: a mini-review. Gerontology 65:323–331. https://doi.org/10. 1159/000492596
- Benitez BA, Cruchaga C (2013) TREM2 and neurodegenerative disease. N Engl J Med 369(16):1567–1568. https://doi.org/10.1056/NEJMc1306509
- Deming Y, Filipello F, Cignarella F, Cantoni C, Hsu S, Mikesell R et al (2019) The MS4A gene cluster is a key modulator of soluble TREM2 and Alzheimer's disease risk. Sci Transl Med 11(505):eaau2291

## **Publisher's Note**

Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

#### Ready to submit your research? Choose BMC and benefit from:

- fast, convenient online submission
- thorough peer review by experienced researchers in your field
- rapid publication on acceptance
- support for research data, including large and complex data types
- gold Open Access which fosters wider collaboration and increased citations
- maximum visibility for your research: over 100M website views per year

#### At BMC, research is always in progress.

Learn more biomedcentral.com/submissions

