**ABCA7 Frameshift Deletion Associated with Alzheimer’s Disease in African Americans**

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Abstract

Objective: To identify a causative variant(s) that may contribute to Alzheimer’s disease (AD) in African Americans (AA) in the ATP-binding cassette, sub-family A (ABC1), member 7 (ABCA7) gene, a known risk factor for late-onset AD.

Methods: Custom capture sequencing was performed on ~150 kb encompassing ABCA7 in 40 AA cases and 37 AA controls carrying the AA risk allele (rs115550680). Association testing was performed for an ABCA7 deletion identified in large AA datasets (discovery n=1,068; replication n=1,749) and whole exome sequencing Caribbean Hispanic (CH) AD families.

Results: A 44 base pair deletion (rs142076058) was identified in all 77 risk genotype carriers, showing that the deletion is in high linkage disequilibrium with the risk allele. The deletion was assessed in a large dataset (531 cases, 527 controls) and, following adjustments for age, sex, and APOE status, was significantly associated with disease (p=0.0002, OR=2.13 [95% CI:1.42-3.20]). An independent dataset replicated the association (447 cases, 880 controls, p=0.0117, OR=1.65 [95% CI:1.12-2.44]), and joint analysis increased the significance (p=1.414x10^-5, OR=1.81 [95% CI:1.38-2.37]). The deletion is common in AA cases (15.2%) and AA controls (9.74%), but in only 0.12% of our NHW cohort. Whole exome sequencing of multiplex, Caribbean Hispanic families identified the deletion co-segregating with disease in a large sibship. The deleted allele produces a stable, detectable RNA strand and is predicted to result in a frameshift mutation (p.Arg578Alafs) that could interfere with protein function.

Conclusion: This common ABCA7 deletion could represent an ethnic specific, pathogenic alteration in Alzheimer’s disease.
Introduction

Alzheimer's disease (AD) is the leading cause of dementia in the elderly. AD occurs at a higher frequency in minority populations, with estimates of AD being twice as frequent in African Americans (AA) compared to non-Hispanic white (NHW) populations. Apolipoprotein E (APOE) was the first gene associated with AD and the ε4 allele confers an increased risk across populations. While APOE ε4 occurs more frequently in AA than NHW, paradoxically, it has a lower effect size in AA. Therefore, while differing ethnicities share risk genes and alleles, the consequences may be different in distinct populations.

Recent studies have identified >20 additional loci associated with late-onset AD, including ABCA7. Although ABCA7 was first implicated in NHW, a genome-wide significant signal was also detected in AA individuals at rs115550680, a position in linkage disequilibrium with the NHW genome wide association study (GWAS) hits. The AA allele confers a higher risk (p=2.21×10^-9, OR=1.79 [95% CI, 1.47-2.12]) than the most significantly associated alleles in NHW. The effect size of the AA ABCA7 allele is comparable to APOE ε4 in AA (p=5.5×10^-47, OR=2.31 [95% CI, 2.19-2.42]). To date, there is no evidence of a functional consequence of the AA ABCA7 risk allele. Therefore, targeted sequencing of ABCA7 was performed to identify potential causative variants. A frameshift deletion was found associated with AD in AA, but virtually absent in NHW. Thus, this deletion potentially represents a common, ethnic-specific, and likely pathogenic alteration that confers risk to Alzheimer's disease.

Methods

Standard Protocol Approvals, Registrations, and Patient Consents
All of the individuals ascertained for this study provided written informed consent prior to their inclusion. If a study participant was not competent to provide consent, immediate next of kin or a legal representative provided written consent on their behalf. All participants were ascertained via a protocol which was approved by the appropriate Institutional Review Board. Oversight of this study falls under the University of Miami IRB #20070307.

Sample Collection

African Americans

Individuals were ascertained for this study following informed consent at the John P. Hussman Institute for Human Genomics (HIHG) at the University of Miami Miller School of Medicine (Miami, FL), North Carolina A&T State University (Greensboro, NC), and Case Western Reserve University (Cleveland, OH) for the HIHG dataset. Each of the participants were ascertained using the protocol approved by the proper Institutional Review Boards. Patients were collected for this study over the course of 10 years, with IRB protocols and amendments being approved at each stage. For the HIHG cohort (discovery), 539 cases were ascertained (415 women and 124 men, mean age of onset 74.0 years [SD 8.5]) as well as 529 controls (403 women and 126 men, mean age at exam 73.1 years [SD 5.4]). The complete HIHG case-control AA cohort (n=1,068) included 47 relatives, giving 1021 independent (unrelated) individuals available for analysis.

Samples from the Alzheimer’s Disease Genetics Consortium (ADGC) were collected as previously described 11. For the ADGC cohort (replication), 687 unrelated cases were ascertained (499 women and 188 men, mean age of onset 78.7 years [SD 8.5]) as well as 1,062 unrelated controls (774 women and 288 men, mean age at exam 78.6 year [SD 6.7]). This subset of the ADGC cohort was independent from the HIHG cohort.
For both HIHG and ADGC datasets, participants underwent rigorous phenotyping and diagnostic criteria following that of the National Institute of Neurological and Communicative Disorders and Stroke – Alzheimer’s Disease and Related Disorders Association, as previously described \(^{11,15,16}\). Furthermore, the cognitive status of controls was measured with either the Mini-Mental State Exam (MMSE) \(^{17}\) or the Modified Mini-Mental State (3MS) \(^{18}\), as well as the Clinical Dementia Rating Scale (CDR) which assesses functional decline \(^{19}\). All of the individuals in both cohorts enrolled self-identified as African American. These data were confirmed by analysis of existing GWAS data \(^{11}\).

**Caribbean Hispanics**

Nineteen multiplex Caribbean Hispanic families initially recruited as a part of the Genetic Epidemiology of Alzheimer’s Disease In Hispanics (EFIGA) family study at Columbia University were utilized. A total of 49 cases and 8 unaffected relatives were involved in this study. Information about patient recruitment, demographics and clinical phenotyping have been previously published \(^{20}\). Each family has at least one member with early-onset AD (age at onset <65 years old).

**Custom Capture and Whole Exome Sequencing**

Custom sequence capture was performed on 77 HIHG samples of African American ancestry (40 cases and 37 controls) all with the AA risk allele. Probes were selected using the Agilent SureDesign program across the region (chr19:983277-1133190, ~150 kb, Agilent Technologies, Santa Clara, CA). 14,636 probes were chosen at a 3x density with the moderately stringent masking setting to cover 84.8% of the region. For whole exome sequencing, samples from the 19 Caribbean Hispanic families were utilized (46 cases and
6 unaffected relatives). Capture and sequence library construction was performed on a Sciclone G3 NGS Workstation (Caliper Life Sciences, PerkinElmer, Waltham, MA) using the SureSelect Human All Exon 50Mb Kit (Agilent Technologies) and the Paired-End Multiplexed Sequencing library kit (Illumina, San Diego, CA) for sequence library preparation. All samples were run on the Illumina HiSeq 2000 and paired end 2x100 sequencing was performed. The sequencing data were processed using the Illumina Real Time Analysis (RTA) base calling pipeline version 1.8. The Burrows-Wheeler Aligner (BWA) was used to map sequences to the hg19 human reference genome and variant calling was performed with the Genome Analysis Toolkit (GATK \cite{21,22}). GATK parameters included base quality score recalibration and duplicate removal \cite{23}. The data were evaluated for deletions and insertions by alignment with Bowtie2 and analysis using the Pindel program \cite{24,25}.

**Sanger Sequencing**

Both the \textit{ABCA7} deletion (rs142076058) and the AA \textit{ABCA7} risk allele (rs115550680) were sequenced using traditional Sanger sequencing. Custom primers were designed with the Primer3 v4.0 program (http://fokker.wi.mit.edu/primer3/input.htm). For the deletion, primers were selected to flank the 44 base pair (bp) deletion in order to perform Sanger sequencing for validation (Deletion-F: AAATCTTCCCGCTTGAGAT, Deletion-R: GGAGCTTAGGTGCAGCTC). Polymerase chain reaction (PCR) experiments were set up with 1.5 mM MgCl\textsubscript{2}, 1.6 M betaine, and touchdown PCR performed. PCR experiments resulted in amplicons of either 450 or 406 bp. Sequencing of the AA risk allele was performed with the following primers (rs115550680-F: GCCAATATGGGCAAAACCATC, rs115550680-R: TCCAAACCCCTGTGATGACC) to generate a 245 bp amplicon. PCR
reactions were set up with 2 mM MgCl$_2$ and touchdown PCR performed. Sequencing reactions were performed utilizing the Big Dye Terminator v3.1 (Life Technologies, Carlsbad, CA), reactions were run on a 3730xl DNA Analyzer (Life Technologies), and results evaluated using the Sequencher v4.10.1 program (Gene Codes Corporation, Ann Arbor, MI).

**TaqMan SNP Genotyping Analysis**

Both the $ABCA7$ deletion (rs142076058) and the AA $ABCA7$ risk allele (rs115550680) were evaluated using TaqMan SNP Genotyping Assay (Life Technologies). The $ABCA7$ deletion was evaluated by a custom designed TaqMan SNP Genotyping Assay designed to recognize the presence or absence of the deletion. This assay had to be ordered as a “non-Human Assay” (forward primer: GCCTGGATCTACTCCGTGAC, reverse primer: GAGGCAGCTGAGGAACCA, FAM probe: GAGACGCGGCTGG – identifies when the sequence is deleted, VIC probe: CGCCATGGGGCT – wild type allele). Samples were amplified for 40 cycles and, when amplification was low, an additional 20 cycles was added. The plates were read on the 7900HT Fast Real-Time PCR machine (Applied Biosystems, Foster City, CA) and data analyzed with the SDS v2.4 software.

**RNA Isolation and RT-PCR**

RNA was isolated from blood collected in PAXgene blood RNA tubes (PreAnalytiX, Hombrechtikon, Switzerland) from 10 samples with and 10 samples without the $ABCA7$ deletion and extracted following the manufacturer’s standard protocol. RNA was quantified on the 2100 BioAnalyzer (Agilent Technologies) and required to have a RIN $\geq$6. Complementary DNA (cDNA) libraries were generated using the iScript Reverse
Transcriptase Supermix for RT-qPCR kit (BIO-RAD). PCR primers were designed to amplify the cDNA across the deletion (cDNA-F: TGTTCCTGCGTGTGCTGA, cDNA-R: AGCAGGAAGCTCTGGGTCAC) and the resulting PCR products resolved on a 2% agarose gel. The wild type allele results in an amplicon of 316 bp, while the allele with the deletion produces a 272 bp amplicon.

**Statistical Analysis**

The GENMOD program, as part of the SAS/STAT software, was utilized to perform the association tests under a logistic regression model. Association tests were performed with adjustments for age, sex, *APOE* status, and relatedness between samples (SAS Institute, Cary, NC). Conditional analysis was performed in PLINK. Fisher’s Exact test was used to evaluate the differences in the alleles frequency of the deletion between African and European populations reported in the ExAC database.

**Results**

We selected 40 AA AD cases and 37 AA controls (aged >65 years) carrying the AA risk allele, rs115550680, to perform custom massively parallel sequencing of a ~150 kb region that includes *ABCA7* as well as eight flanking genes and a small nuclear RNA. Samples were sequenced to an average depth of over 1000x and evaluated for single nucleotide variants (SNVs) and insertions and deletions. 1,120 SNVs were detected by sequencing with 11 variants showing different frequencies in cases and controls (p<0.1, Table e-1). In addition, a 44 base pair (bp) deletion (rs142076058, p.Arg578Alafs) located ~3.5 kb upstream of the AA risk allele was identified in all 77 individuals, suggesting that it is in high linkage disequilibrium with the risk allele (Figure 1).
To further evaluate the rs142076058 deletion, a custom TaqMan genotyping assay was designed to evaluate the deletion in our larger AA cohort, designated as HIHG. Following adjustments for age, sex, and APOE status, the deletion was found to be significantly associated with AD (p=0.0002, OR=2.13 [95% CI:1.42-3.20], Table 1) in 531 cases and 527 controls. The deletion occurred in 9.3% of control individuals, but 16.2% of AD cases. A subset of eight individuals were pathologically-confirmed cases of AD; two were found to carry the deletion while the remaining six did not. The AA risk allele was also genotyped in this dataset. The risk allele was significantly associated with AD (p=0.0005, OR=2.07 [95% CI:1.38-3.13]) and in linkage disequilibrium with the deletion (D'=1.000, r²=0.995, Table e-2 and e-3). The top SNPs in ABCA7 previously reported in NHW studies were also found to be in LD with the deletion (Table e-4) 8,9,11. Genotyping was also performed on our NHW AD samples (n=3,275), but only 4 individuals were identified with the deletion (0.12%), all of whom carried the AA risk allele, indicating that the genomic fragment carrying the deletion in these individuals may be of African descent.

An independent AA dataset from the Alzheimer’s Disease Genetics Consortium (ADGC) was evaluated in the same manner and the deletion was again significantly associated with AD (p=0.0117, OR=1.65 [95% CI:1.12-2.44], Table 1), occurring in 10.0% of control individuals and 14.9% of AD cases. Joint analysis of the two cohorts resulted in an overall significant association (p=1.414x10⁻⁵, OR=1.81 [95% CI:1.38-2.37], Table 1). Association testing was also performed for each dataset without APOE adjustment; more significant results were obtained with APOE adjustment, demonstrating that APOE did not influence the association (Table e-5). Examination of the ages of cases and controls with and without the deletion did not find a significant difference between any of these groups (Table e-6). To examine the association with AD in another ethnic group with a high level of African
ancestry (~42%), we evaluated whole exome sequencing data on 19 Caribbean Hispanic families from the Dominican Republic with multiple affected AD subjects. In addition to a relatively high level of African ancestry, Caribbean Hispanics are highly inbred and have a high incidence of AD, and are thus enriched for AD genetic risk factors. We independently identified the same 44 bp deletion from whole exome sequencing of three affected individuals from a large Caribbean Hispanic family. Subsequent examination of the family revealed that the deletion segregated in a large sibship in the family (Figure 2). Both the deletion and AA risk allele were isolated in all 7 siblings who clinically presented in a range from Alzheimer’s disease (individuals 5, 6, 8, 12, and 99) to milder stages of dementia (individuals 7 and 11). Haplotype analysis around the ABCA7 deletion using SNP data in the 1MB flanking region on the family revealed that an affected aunt who does not carry the deletion, individual 3 (Figure 2), has distinct ancestral haplotypes from the family members in the large sibship with the deletion (Table e-7). This finding suggests that individual 3’s AD phenotype can be attributed to other genetic factors and that the ABCA7 deletion is highly penetrant in the sibship. Since several members of this CH family were known to have early onset AD (age at onset <65 years), we examined the entire AA AD cohort (both HIHG and ADGC) to determine if there was an effect of the deletion on age of onset in AD. We did not find a significant difference in the age of onset in cases with the deletion (75.6 years [SD 9.6]) compared to cases absent for the deletion (76.8 years [SD 8.7], p=0.09).

To determine whether the ABCA7 allele with the deletion was being transcribed, RNA was isolated from the blood of AA individuals both with and without the ABCA7 deletion. Reverse transcription-PCR across the deletion region demonstrated that the allele carrying the deletion is transcribed and produces a stable, detectable RNA strand (Figure 3).
This deletion was previously reported in the Exome Aggregation Consortium (ExAC: http://exac.broadinstitute.org) [6, 2015], a repository of 60,706 unrelated individuals from six distinct ethnic groups. In the ExAC dataset, the deletion was found in 7.77% of individuals of African ancestry and 0.95% of Latino individuals, but was absent from individuals of European ancestry. This difference in population frequencies between the African and European populations was highly significant (p<1x10^{-10}).

**Discussion**

We identified a 44 bp deletion in *ABCA7* that is significantly associated with AD in individuals of African ancestry. While the deletion did occur in unaffected individuals, it was found at a higher frequency in individuals with AD (15.2% of cases versus 9.74% of controls), implicating it as a risk factor for disease. This reaffirms that the deletion is likely to be of African ancestry. Furthermore, the combined cases from the HIHG and ADGC datasets had the deletion at a frequency of 15.2%, approximately twice as high as that identified with through ExAC African populations (7.8%), lending additional evidence of a relationship to disease. The deletion was also independently identified in an AD family from the Dominican Republic, a population that has a relatively high level of African ancestry, 41.8% [27]. Examination of the linkage disequilibrium of the deletion with the top three previously reported SNPs found a high D’ across all locations, but only a significant r^2 with the African specific risk allele (Table e-4), further supporting that distinct alleles confer AD risk in different ethnicities [8,9,11].

The deletion is predicted to cause a frameshift at amino acid 578, encoding for 168 incorrect amino acids before stopping prematurely compared to the largest isoform that generates a protein of 2146 amino acids (Figure 1B). Since we were able to detect RNA
from the allele with the deletion, it is possible that this RNA generates an aberrant protein that interferes with the wild type 2,146 amino acid protein. Within the first 578 amino acids, two transmembrane regions are conserved and would be maintained by the mutated protein (Figure 1B). However, both AAA domains and 9 additional transmembrane domains would be predicted to be lost in this truncated protein, and thereby interfere with the protein’s function of exporting the lipid phosphatidylserine 29. Alternatively, the shortened transcript may be subjected to nonsense-mediated decay (NMD), as was seen in the Glu709fs alteration identified in NHW 30. While some loss-of-function variants in ABCA7 were identified in NHW populations that may contribute to AD pathogenicity, these are rare variants and may only partially contribute to the NHW GWAS signal 30-32. A few previously reported loss-of-function variants have demonstrated a functional consequence including the Glu709fs variant undergoing NMD and the c.5570+5G>C alteration led to aberrant splicing 30,31 (Figure 1B). Therefore, this study may be the first to connect a potentially pathogenic and common alteration with a GWAS signal in ABCA7.

ABCA7 is a member of the ATP-binding cassette (ABC) transporter family, a large group of 49 genes that encodes for membrane proteines that facilitate the movement of substrates across cell membranes 33,34. ABCA7 is expressed in the brain in neurons and microglia 35,36. There is evidence both in patients and animal models demonstrating that inadequate levels of ABCA7 may be directly correlated with Alzheimer pathogenesis 37-39. The ABCA7 protein is involved in the processing of amyloid precursor protein (APP 40). In addition, evidence has shown that ABCA7 acts in the phagocytic pathway via extracellular signal-regulated kinase signaling e-1,e-2. ABCA7 is not the only ABC transporter gene linked to AD; ABCA1, ABCB1, ABCC1, ABCG1, ABCG2, and ABCG4 are all implicated in Aβ regulation e-3,e-8. Furthermore, a study identified rare loss-of-function alterations in NHW patients
diagnosed with Parkinson’s disease, including specific variants previously reported in AD individuals, demonstrating that this gene may contribute to the risk of multiple neurodegenerative disorders \(^{e-9}\).

Therefore, the results of this study demonstrate that there is a 44 base pair deletion in \(ABCA7\) that is significantly associated with AD and in linkage disequilibrium with the previously identified AA risk allele. The deletion was relatively frequent in our large AA AD cohorts, independently identified in 1 of 19 Caribbean Hispanic AD families, and virtually absent from our large NHW AD cohort. Thus, the deletion could represent a common, ethnic-specific alteration that confers risk of Alzheimer’s disease in populations with African ancestry.
**Figure Legends**

**Figure 1. Location of the Deletion in the ABCA7 gene and protein.** A. The ABCA7 gene (chr19:1,040,103-1,065,571, hg38), the 44 base pair deletion (blue), the AA risk allele (blue and underlined\(^{11}\)), and three NHW risk alleles (rs3764650\(^8\), rs3752246\(^9\), rs4147929\(^{10}\)). B. The wild type ABCA7 protein (2146 amino acids) and the location of frameshift deletion (blue) identified in this study. Below, the protein the predicted to be generated from deletion would contain only two of the eleven transmembrane domains (yellow) and neither of the two AAA domains (green), but incorporate 168 aberrant amino acids (black). The remaining frameshift, nonsense, and splicing variants designated are rare alterations (<1 % MAF) previously reported in NHW populations to be associated with AD\(^{33-35}\).

**Figure 2. Pedigree of an AD family from the Dominican Republic with the ABCA7 Deletion.** Family 360 has six individuals diagnosed with Alzheimer’s disease, as well as two individuals presenting with mild dementia. The numbers beneath each individual in the pedigree represent the individual’s sample number, the age of onset of Alzheimer’s disease (for AD cases) or the age at exam, and the APOE genotype. Seven siblings all carry the ABCA7 deletion as well as the AA risk allele, while individual 3 did not have either the deletion or the AA risk allele.

**Figure 3. Deletion Allele Produces an RNA Transcript.** A. RT-PCR from cDNA of three samples without the ABCA7 deletion (+/+), and three samples heterozygous for the ABCA7 deletion (deletion/+). All samples produce an amplicon of 316 base pairs, but only the samples with deletion generate a lower, 272 base pair amplicon (arrow). B. Sanger sequencing from the 5’ end of the deletion in an AA control lacking the deletion and AD-
specific line heterozygous for the deletion. The arrow denotes where the deleted allele begins to be out of frame with the wild type allele C. Sanger sequencing from the 3’ end of the deletion from the same control and AD individuals.
**Table 1.** Association Testing of the Deletion in AA Cohorts

|               | samples with deletion/ total samples (%) | odds ratio | 95% CI     | Pr > |Z| |
|---------------|------------------------------------------|------------|------------|------|---|
| **HIHG<sup>a</sup>** |                                          |            |            |      |   |
| cases         | 86/531 (16.2)                            | 2.13       | 1.42-3.20  | 0.0002 |   |
| controls      | 49/527 (9.3)                             |            |            |       |   |
| **ADGC<sup>b</sup>** |                                          |            |            |      |   |
| cases         | 63/447 (14.9)                            | 1.65       | 1.12-2.44  | 0.0117 |   |
| controls      | 88/880 (10.0)                            |            |            |       |   |
| **Joint analysis** |                                          |            |            |      |   |
| cases         | 149/978 (15.2)                           | 1.81       | 1.38-2.37  | 1.414x10^-5 | |
| controls      | 137/1407 (9.7)                           |            |            |       |   |

<sup>a</sup>HIHG – John P. Hussman Institute for Human Genomics, University of Miami

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References


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