Genomic Convergence to Identify Candidate Genes for Alzheimer Disease on Chromosome 10

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ABSTRACT: A broad region of chromosome 10 (chr10) has engendered continued interest in the etiology of late-onset Alzheimer Disease (LOAD) from both linkage and candidate gene studies. However, there is a very extensive heterogeneity on chr10. We converged linkage analysis and gene expression data using the concept of genomic convergence that suggests that genes showing positive results across multiple different data types are more likely to be involved in AD. We identified and examined 28 genes on chr10 for association with AD in a Caucasian case–control dataset of 506 cases and 558 controls with substantial clinical information. The cases were all LOAD (minimum age at onset ≥60 years). Both single marker and haplotypic associations were tested in the overall dataset and eight subsets defined by age, gender, ApoE, and clinical status. PTPLA showed allelic, genotypic, and haplotypic association in the overall dataset. SORCS1 was significant in the female subset (allelic association p = 0.0025) and most significant in the female subset (allelic association p = 0.00002, a three-locus haplotype had p = 0.00055). Odds ratio of SORCS1 in the female subset was 1.7 (p < 0.0001). SORCS1 is an interesting candidate gene involved in the AD pathway. Therefore, genetic variations in PTPLA and SORCS1 may be associated and have modest effect to the risk of AD by affecting AD pathway. The replication of the effect of these genes in different study populations and search for susceptible variants and functional studies of these genes are necessary to get a better understanding of the roles of the genes in Alzheimer disease. 


KEY WORDS: Alzheimer disease; late-onset Alzheimer disease; LOAD; genomic convergence; association; candidate genes; PTPLA; SORCS1

Introduction

Alzheimer disease (AD; MIM# 104300) is the leading cause of dementia in the elderly and the most common form of dementia occurring after the age of 40 [Rocca et al., 1991; Schoenberg et al., 1987]. It is a devastating neurodegenerative disorder of later life with a largely unknown but complex etiology that includes a strong genetic component [Growden, 1995; Khachaturian, 1985]. Understanding the genetics of AD is critical to developing new treatments.

Genetic studies have identified three genes underlying early onset AD. This form can be caused by over 120 mutations in genes encoding β-amyloid precursor protein (APP; MIM# 104760), presenilin 1 (PS1; MIM# 104311), and presenilin 2 (PS2; MIM# 600759) [Goate et al., 1991; Levy-Lahad et al., 1995; Rocchi et al., 2003; Sherrington et al., 1995]. To date, five late-onset AD genome-wide linkage screens have been reported [Blacker et al., 2003; Kehoe et al., 1999; Myers et al., 2002; Pericak-Vance et al., 1997, 1998, 2000]. In addition, numerous studies have tested hundreds of candidate genes for involvement in AD [Bertram et al., 2007]. However, no single method can give us an answer in dissecting complex disease traits. A promising solution to the growing data flood from the latest genomic technologies (e.g., genome-wide association) is genomic convergence. Genomic convergence is a multifactor approach used in genetic research that combines different data analysis results from independent types to identify and prioritize susceptibility genes for a complex disease [Hauser et al., 2003]. Examples include genetic linkage data, association data, gene expression data, and biological function.

Our previous linkage and candidate gene studies suggested a more extensive heterogeneous genetic background on chromosome 10 for AD than we previously appreciated [Liang et al., 2007]. Rare and common polymorphisms in multiple genes may be involved together with nongenetic factors in this complex disease. Therefore, combining different sources of data is important to selecting and prioritizing candidate genes.

In a recent study [Xu et al., 2007] gene expression levels were compared in the brain tissue from AD patients and controls using the Serial Analysis of Gene Expression (SAGE) method. SAGE is a powerful method for profiling transcripts expressed in a given tissue [Velculescu et al., 1995], and provides expression data for thousands of gene products without a priori knowledge of their function. Brain samples from AD cases (ApoE 4/4, ApoE 3/4, and ApoE 3/3 genotypes) and controls (ApoE 3/3 genotypes) were
chosen to establish SAGE libraries to analyze the differential gene expression between AD cases and controls as described elsewhere [Xu et al., 2007]. We converged the previous genetic linkage screen results with the SAGE gene expression data to identify a set of 28 genes that reside under the linkage peak and are differentially expressed between patients and controls. Numerous SNPs in these genes were genotyped and examined for association with AD.

Common diseases are likely to have complex etiology. Multiple genes with multifaceted interactions among genes and other risk factors undoubtedly play roles in the AD process. Multifactor dimensionality reduction (MDR) is a computational data reduction method that can identify gene–gene interactions without requiring very large sample sizes. Because MDR is a nonparametric, genetic model-free method, no hypothesis concerning any statistical parameter and no genetic inheritance model is assumed [Ritchie et al., 2001]. In the present study, we applied MDR to the 28 candidate genes to test for potential gene–gene interactions involved in AD.

Materials and Methods

Study Populations

The case–control data set consisted of 1,064 individuals (506 cases and 558 controls; Table 1) with substantial clinical information. The samples were collected by the Center for Human Genetics Research (CHGR) at Vanderbilt University and the Miami Institute for Human Genomics (MIHG) at the University of Miami. AD was diagnosed according to the NINCDS-ADRDA criteria [McKhann et al., 1984]. The cases were all late onset AD (minimum age at onset (AAO) ≥ 60 years). There were no significant differences in age at exam or gender between cases and controls (Table 1). All controls were ascertained in the same catchment area as cases, and had results within the normal range in the Mini-Mental State Exam (MMSE) or Modified Mini-Mental State Exam (3MS). To test the potential for heterogeneity effects in the data set, the overall data set was subsetted according to four variables: gender, ApoE status, age of onset, and clinical status. The distribution is shown in the Supplemental Table S1. Written consent was obtained from all participants in agreement with protocols approved by the institutional review board at each contributing center.

We also used independent family-based data sets as the validation sets to confirm any significant association identified in the case–control data set (Table 2). All individuals included in this study were Caucasian late-onset AD (LOAD) patients. AD diagnosis criteria were same as those for the case–control data set.

Samples were ascertained by the following centers: the NCRAD repository at Indiana University (NCRAD); the Collaborative Alzheimer Project (CAP), including the University of Miami and Vanderbilt University and University of California at Los Angeles; and the National Institute of Mental Health repository (NIMH).

Following informed consent, blood samples were collected from each individual. Genomic DNA was extracted from whole blood by use of the Puregene system (Gentra Systems, Minneapolis, MN). All samples were coded and stored at 4°C until used.

Gene Selection

The SAGE method was used to compare the gene expression levels in the brain tissue from AD patients and controls as described elsewhere [Li et al., 2006; Velculescu et al., 1995]. Briefly, six SAGE libraries were generated using hippocampus collected in the Kathleen Price Bryan Brain Bank, at the Duke University Alzheimer Disease Research Center, and the Brain Bank of the Center for Human Genetics (CHG), Duke University Medical Center, following a rapid autopsy protocol [Hulette et al., 1997]. Four libraries were based on the short tag (14 bp, AD ApoE 4/4, ApoE 3/4, ApoE 3/3, and control ApoE 3/3). Two libraries were based on the long tag (21–22 bp, AD ApoE 3/3, and control ApoE 3/3). The number of times a tag observed in tissue was extracted from each library and compared between the AD and control samples using SAGE software to form a compared ShortSAGE database. Chi-square and Fisher exact tests, as previously described [Hauser et al., 2003], were used to test differences in expression levels between AD and control for each tag in each compared SAGE data set. The SAGE tags were compared to the UniGene to interpret the SAGE results. The false discovery rate (FDR) was applied to the SAGE data to correct for multiple comparisons. $p<10^{-10}$ was used as the cutoff to select genes whose expression levels were highly significantly different between AD cases and controls. Then the previous linkage study results for chromosome 10, shown in Table 3, were converted with gene expression data so that only differentially expressed genes that were also under linkage peaks in previous linkage screens were selected for the next step association analysis.

Table 1. Alzheimer Disease Case Control Data Set

<table>
<thead>
<tr>
<th>Sample size (N)</th>
<th>Case</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>311</td>
<td>337</td>
</tr>
<tr>
<td>Male</td>
<td>195</td>
<td>221</td>
</tr>
<tr>
<td>Age of onset</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All</td>
<td>72.5</td>
<td>72.8</td>
</tr>
<tr>
<td>Female</td>
<td>72.1</td>
<td>75.9</td>
</tr>
<tr>
<td>Male</td>
<td>75.9</td>
<td>74.3</td>
</tr>
<tr>
<td>Age of exam</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All</td>
<td>75.9</td>
<td>73.5</td>
</tr>
<tr>
<td>Female</td>
<td>76.2</td>
<td>75.5</td>
</tr>
<tr>
<td>Male</td>
<td>75.8</td>
<td>74.3</td>
</tr>
</tbody>
</table>

Table 2. Family-Based Data Set for Alzheimer Disease

<table>
<thead>
<tr>
<th>Family</th>
<th>Overall</th>
<th>NIMH</th>
<th>NCRAD</th>
<th>CAP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total pedigrees</td>
<td>730</td>
<td>352</td>
<td>154</td>
<td>224</td>
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<tr>
<td>Affected individuals</td>
<td>1,521</td>
<td>807</td>
<td>315</td>
<td>390</td>
</tr>
<tr>
<td>Unaffected individuals</td>
<td>974</td>
<td>331</td>
<td>162</td>
<td>481</td>
</tr>
<tr>
<td>Discordant Sib Pairs (DSP)</td>
<td>1,337</td>
<td>629</td>
<td>269</td>
<td>439</td>
</tr>
<tr>
<td>Independent Discordant Sib Pairs</td>
<td>674</td>
<td>283</td>
<td>129</td>
<td>262</td>
</tr>
<tr>
<td>Pedigrees with at least one DSP</td>
<td>406</td>
<td>165</td>
<td>75</td>
<td>166</td>
</tr>
<tr>
<td>Pedigrees with at least one ARP</td>
<td>188</td>
<td>66</td>
<td>26</td>
<td>96</td>
</tr>
<tr>
<td>Pedigrees with at least one ARP</td>
<td>64</td>
<td>31</td>
<td>11</td>
<td>22</td>
</tr>
</tbody>
</table>
prioritizes these SNPs on their tagging for linkage disequilibrium based on the LD bin algorithm in LD select [Carlson et al., 2004] using HapMap data, SNP allele frequencies and source, function, regulatory potential, and repeat status. Because we used the Illumina Goldengate genotyping method, we only selected SNPs with Illumina SNPscore greater than 0.6 to ensure the genotyping quality. An MAF greater than 20% was used as the cutoff for selecting the intronic SNPs. SNPs in conserved regions of eight-gene alignment (Human/chimp/mouse/rat/dog/chicken/fugu/zebra_fish, based on UCSC database), conserved transcription factor binding sites (TFBSs) in the human/mouse/rat alignment, CpG island, and microRNA genes were selected if the MAFs were greater than 10%. Coding SNPs were selected when the MAF was greater than 1%. To further reduce the SNPs to a manageable number, there were selected at the one SNP/10 kb spacing. Because TNFrsF6 showed significant results in linkage, candidate gene studies, and SAGE gene expression analysis [Bertram et al., 2000; Feuk et al., 2000], more weight was given to it by genotyping one SNP per five kilobases.

The final set of 667 SNPs on chr10 was genotyped using the Illumina GoldenGate Oligo Pool assay (OPA) on an Illumina BeadStation 500 GX (Illumina Inc., San Diego, CA). DNA samples from cases and controls were randomly sorted, and duplicated samples were implemented across plates for genotyping quality control.

We tested all SNPs for consistency with HWE in Haploview program [Barrett et al., 2005]. SNPs were excluded if HWE p < 0.001. TagSNPs were selected at different linkage disequilibrium levels using the Tagger option incorporated in Haploview for the future haplotypic association tests and gene–gene interaction tests.

### Statistical Methods

#### Association analysis in case–control data set

In the case–control data set, allelic association for single SNPs was tested using the χ² test in Haploview [Barrett et al., 2005]. Genotypic association was assessed by a 2 × 3 contingency table likelihood ratio test in SAS program (SAS Institute, Cary, NC, 2003). Haplotypic association was analyzed using haplo.stats [Schaid et al., 2002]. To save calculation time and satisfy computer memory limits, small genes (with less than eight genotyped SNPs in the gene) were analyzed using individuals with less than 50% missing genotypes. Genes of medium size (with 8 to 26 genotyped SNPs) were analyzed using only individuals who had complete genotypes on all analyzed loci in that gene. There are seven large genes that were genotyped for 38 to 124 SNPs. They were analyzed on individuals without any missing genotypes and only tagSNPs with r² < 0.5 were used. In addition, the tagSNPs were separated into groups of 10 with 2 SNPs overlapping across the groups. We estimated haplotype frequencies and tested association of each haplotype with a frequency of at least 1% in our case–control data set with age and gender-adjusted score statistics. Haplotype logistic regression was modeled using a GLM algorithm including age and gender as covariates. The most frequent haplotype occurring in the similar percentage of cases and controls was selected as the referent haplotype. To evaluate the association of subsets of alleles from the full haplotype, a sliding window of three SNPs was used and the global score statistics were calculated.

Conditional analyses were conducted by dividing the case–control data set by known risk factors including age, gender, and ApoE status. The difference in allele frequency between cases and controls was tested by a χ² test or Fisher’s exact test when applicable (e.g., cell size < 5). We also applied logistic regression to estimate the effects of the SNPs showing the most significant results across different analyses after controlling for age, gender, and ApoE status. Nominal significance was declared at p = 0.05 and FDR was used to correct multiple testing. Haploview and SAS (SAS institute) programs were used for the analyses.

### Association analysis in family data set

Family based association analysis was used to follow up the SNP with the strongest association in the female subset. The allelic association analyses were conducted using the association in the presence of linkage (APL) analysis [Martin et al., 2003b] and the pedigree disequilibrium test (PDT) [Martin et al., 2000]. APL and PDT each have distinct advantages. APL can correctly infer missing parental genotypes in regions of linkage by estimating identity-by-descent (IBD) parameters and is a more powerful test under those conditions than PDT. However, APL only uses nuclear families for the association test. PDT can use data from extended pedigrees and gains power substantially when extended pedigree data are available. Genotype-PDT (GenoPDT) is an extension of PDT and was used to assess association between genotypes and risk of AD in the family data [Martin et al., 2003a].

### Test for gene–gene interaction

TagSNPs were selected using r² > 0.5 as a threshold to exclude SNPs from the interaction analysis that were in high LD by the Tagger option in Haploview. Potential gene–gene interactions were identified using MDR. MDR performs an exhaustive search of all possible single-locus through k-locus interactions to evaluate all possible high/low risk models of disease [Ritchie and Motsinger, 2005]. The statistical significance of the final best model was determined through permutation testing with 1,000 permuted data sets. The significance of the final model was determined by comparing the prediction error and crossvalidation consistency of the final model with the distribution. A p value was extracted for the model by its theoretical location in the permutation distribution [Motsinger et al., 2006]. In considering the possible unbalanced ratio of cases to controls in the data set, a balanced accuracy approach was used to evaluate the performance of the models [Velez et al., 2007].

In the present study, one, two, and three locus combinations were run and logistic regression (SAS program) was used to verify the interaction between genes in the identified model.
Results

Genomic Convergence Identified Genes and Genotyped SNPs

SAGE analysis revealed 41 genes on chromosome 10 that were differentially expressed between AD cases and controls with \( p < 10^{-10} \). Twenty-eight of those genes were also under previously identified linkage peaks [Bertram et al., 2000; Liang et al., 2007; Myers et al., 2002; Pericak-Vance et al., 2000; Zuebenko et al., 1998], and thus considered for detailed analysis. The genomic size of the genes ranged from 10 kb to 1.3 Mb. The number of exons ranged from 2 to 38. Depending on the size of the gene, we genotyped 2–124 SNPs in the genes to for an average spacing of one SNP/10 kb. The gene description, size, and number of genotyped SNPs were summarized in the Supplemental Table S2.

Of the 667 genotyped SNPs 70 SNPs were dropped from the analysis because of monomorphism, being out of HWE, or assay problems. Specifically, 58 SNPs failed Illumina quality control criteria, 10 SNPs were monomorphic and two SNPs were out of HWE in controls at \( p < 10^{-5} \) (Bonferroni correction for 667 markers at \( p = 0.05 \)). In total, there were 597 SNPs that remained in the final analyses. Eighty percent of those were intronic SNPs, 17% were functional SNPs (exon, promoter, UTR, exon/intron boundary, etc.) and the other 3% were from the 5’-upstream and 3’-downstream regions (Supplemental Fig. S1).

Association Analysis

The allelic association of 597 genotyped SNPs was analyzed on the overall data set and eight subsets. Twenty-four SNPs in eight genes (PTPLA; MIM* 610467, SORCS1; MIM* 606283, PRKG1; MIM* 176894, SVIL; MIM* 604126, ACTRI1A; MIM* 605143, BA108L7.2, CAMK1D; MIM* 607957, and PDZK7 [approved symbol PDZD7]) showed association with AD at nominal \( p < 0.05 \) (Fig. 1). After we applied FDR (\( q = 0.20 \)) based on the number of genotyped SNPs in each gene to correct for multiple testing, two SNPs in two genes remained significant. The first SNP was rs10508533 in PTPLA with \( p = 0.0022 \); the second is rs17277986 in SORCS1 gene with \( p = 0.0025 \) (Fig. 1). There were additional SNPs showing significant results in each subset defined by age, gender, ApoE status, and clinical status. Five out of eight associated genes in the overall data set showed association in three or more subsets (Supplemental Table S3). These five genes were PTPLA, SORCS1, SVIL, PRKG1, and PDZK7 (approved symbol PDZD7). The female subset had the most significant result (Supplemental Table S3): \( p = 0.00002 \) for a SNP (rs17277986) in SORCS1 (Fig. 2). Close to rs17277986, four tag SNPs (rs2900717, rs10884399, rs11193170, rs4918280) were also significant at \( p < 0.005 \). rs17277986 remained significant after FDR correction for multiple testing. The odds ratio (OR) of having the risk allele versus not having the risk allele was 1.34 (95% confidence interval [CI] = 1.11–1.63, \( p = 0.003 \)) in the overall data set. In the female subset, the OR was 1.70 with 95% CI of 1.33–2.19, and the p value was very significant (\( p < 0.0001 \)). The OR was also significant after the adjustment for age, gender, and ApoE status (data not shown). SNP rs6480499 in PRKG1 was also significant in the female subset after FDR correction (Fig. 2).

Twenty-seven SNPs in 12 genes (CACNB2; MIM* 600003, PTPLA, SORCS1, SVIL, SORBS1; MIM* 605264, PRKG1, CAMK1D, CNNM2; MIM* 607803, PDZK7 [approved symbol PDZD7], TCF7L2; MIM* 600228, TNFRSF6; MIM* 134637, and NT5C2; MIM* 600417) showed association with AD at nominal \( p < 0.05 \) in the genotypic analysis (Supplemental Fig. S2). SNPs in

Analytical Convergence of the Genetic Association Tests

Across all the association analyses including allelic, genotypic, and haplotypic association tests, several genes showed association. After we applied FDR to correct for multiple testing for allelic and genotypic association tests, only five genes showed association in
at least one of the analyses. SNPs in PTPLA were significant in the overall data set across all three association tests. SNPs in SORCS1 had a significant allelic effect in the overall data set, and it was highly significant in the female subset across all three association tests. Although SNPs in CACNB2 did not show allelic association in the overall data set, SNP rs1277738 in CACNB2 had the strongest genotypic effect in overall data set, and showed genotypic association at nominal \( p < 0.05 \) in four subsets (age-of-onset between 60 and 75, ApoE-negative, female, and possible AD). SNPs in SVIL were marginally significant in the haplotypic association test in the overall data set.

**Follow-up of SNPs in SORCS1 and PTPLA Genes**

To confirm the effect of SORCS1, we genotyped the most significant SNP (rs17277986) in our validation data sets. When we analyzed female and male individuals together, none of the datasets showed significant results at \( p < 0.05 \). When we analyzed females only, the overall data set was not significant (\( p = 0.06 \)) (Table 4). The results in family data sets were similar to the population based case–controls data set, but not significant after correction for multiple comparisons.

The two SNPs genotyped in PTPLA were significant across all the previous association analyses (rs10508533 had OR = 1.34 with \( p = 0.003 \); rs4453117 had OR = 1.31 with \( p = 0.003 \)). This gene is small, spanning only 27 kb. We followed up the results by genotyping seven more SNPs in the case–control data set to give this gene a better coverage. The linkage disequilibrium between all genotyped SNPs in cases is shown in the Supplemental Figure S4. Most of the SNPs were significant in allelic and genotypic association test (Table 5). The overall haplotypic association test considering all SNPs in the gene was significant with global score statistic \( p = 0.03 \). The three-SNP sliding window haplotypic association test was significant for all haplotypes (Supplemental Fig. S5).

**Gene–Gene Interaction among Genes on Chromosome 10**

**Overall data set**

To reduce the computational burden and to eliminate highly redundant data, 300 tagSNPs were selected to represent the 597 SNPs at an \( r^2 > 0.5 \) threshold to test for gene–gene interaction in MDR (Table 6). The single best model was found between two SNPs in CACNB2. This two-locus model involves rs1277738 and rs10741083. The average prediction accuracy was 51.43%, and the crossvalidation consistency was 40%, using fivefold crossvalidation. Because other models outperformed this model in other crossvalidation intervals, the \( p \) value was not significant in the permutation test (\( p = 0.37 \)). However, this model is in the top 10 models in three out five of the crossvalidation intervals, being the top model 67% of the time.

When we eliminate noise by using the “FORCLOCI” function in MDR and let the program do the permutation only on the two SNPs in the best two-locus model, the permutation \( p \) value was significant (\( p = 0.003 \)). The result was confirmed in logistic regression when including the interaction term. The interaction was significant with \( p = 0.004 \) in the model. Table 6 shows that SNP rs1277738 in CACNB2 was consistently identified across one-, two-, and three-locus models in MDR. It showed a main effect (best one-locus model) in the MDR analysis. This SNP also had strongest genotypic effect in single marker association test (\( p = 0.003 \)).

Because several models may perform almost equally in different cross validation intervals, looking at only the top model might not be sufficient to identify important genes in this complex disease. Therefore, we considered the frequency of each gene in the top ten MDR models ranked by prediction accuracy. Table 7 shows the three most frequently identified genes in the top 10 MDR models. CACNB2 was in the top 10 models among all crossvalidation intervals in the one-, three-locus models, and in 80% of crossvalidations for two-locus model. In more than 60% of crossvalidation intervals, PTPLA was in the top 10 MDR models. SORCS1 was in top 10 models among 40% of crossvalidation intervals.

**Female subset**

Our previous analysis showed a strong effect of SORCS1 in the female subset, so we performed MDR in this subset as well (Table 8). The single SNP rs17277986 in SORCS1 was identified as the single best model with prediction accuracy of 54.74 and 80% crossvalidation consistency (\( p = 0.10 \)). This SNP was the one showing the strongest single marker allelic effect in the female data set. No two-locus model had a prediction accuracy greater than 50%. The three-locus model contains rs17277986 in SORCS1 and two SNPs (rs1409207 and rs2482100) in CACNB2 with 52.49% prediction accuracy and 40% crossvalidation consistency. Although the permutation \( p \) was not significant at \( p < 0.05 \), the SORCS1 gene was always in the top 10 MDR models among all crossvalidation intervals for one-, two-locus models. CACNB2 was in the top 10 one-locus models among all crossvalidation intervals.

**Table 4. Family-Based Association Tests for rs17277986 in SORCS1 in Validation Data Sets**

<table>
<thead>
<tr>
<th></th>
<th>Female and male</th>
<th>Female only</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>All CAP NIMH NCRAD</td>
<td>All CAP NIMH NCRAD</td>
</tr>
<tr>
<td>APL</td>
<td>0.189 0.696 0.301</td>
<td>0.189 0.575 0.309 0.010</td>
</tr>
<tr>
<td>Sum_PDT</td>
<td>0.862 0.085 0.099 0.742</td>
<td>0.904 0.030 0.254 0.668</td>
</tr>
<tr>
<td>Geno_PDT</td>
<td>0.606 0.269 0.052 0.402</td>
<td>0.798 0.103 0.182 0.562</td>
</tr>
</tbody>
</table>

All, combined data set; CAP, the Collaborative Alzheimer Project; NCRAD, the NCRAD repository at Indiana University; NIMH, the National Institute of Mental Health repository.
Given the assumption that genes showing positive association in multiple tests have higher likelihood of being important in the disease process, we performed an analytical convergence on the results from allelic, genotypic, and haplotypic association tests. The most consistently associated SNPs across the three analyses are rs10508533 in PTPLA in the overall data set (allelic association $p = 0.0022$) and rs17277986 in SORCS1 in the female subset (allelic association $p = 0.00002$).

PTPLA is protein tyrosine phosphatase-like, member A. Little is known about its function, but it could be involved in the protein phosphorylation process that is important for the function of some proteins in the AD pathology. Among all nine genotyped SNPs in PTPLA, seven showed allelic association and six showed genotypic association. Three of those six SNPs are functional SNPs. This gene is not in the same LD block with any nearby genes based on HapMap data. Therefore, the three functional SNPs could have potential roles in the biological function of PTPLA gene or are in LD with a susceptible SNP affecting PTPLA function, but are unlikely to be in LD with susceptibility SNPs in another gene.

SORCS1 encodes sortilin-related VPS10p domain containing type 1 receptor. It contains a leucine-rich domain and mediates intracellular sorting and trafficking functions. It is highly expressed in the brain [Hermey et al., 1999] and neuronal activity can differentially affect its expression [Hermey et al., 2004]. SORCS1 is a substrate of γ-secretase and γ-secretase cuts amyloid precursor protein (APP) and generates amyloid β peptide (Aβ), one of the hallmarks of AD. rs17277986 in SORCS1 was significant in the overall, age-of-onset greater than 76 years old, ApoE 4-positive, and definite/probAD data sets, with $p = 0.0025, 0.0178, 0.0073,$ and $0.0034$ respectively. It was most significant in the female subset ($p = 0.00002$). It survived not only an FDR multiple testing correction, but also a more conservative Bonferroni correction. Four nearby tagSNPs ($r^2<0.8$) were also significant in the female subset ($p = 0.0008, 0.002, 0.0064, 0.0018$ for rs2900717, rs10884399, rs11193170, and rs4918280, respectively). Another SNP (rs12571141) had a $p = 0.00006$ in this subset, but it was in high LD with rs17277986 ($r^2 = 0.97$), suggesting it was measuring the same effect. Haplotypes containing the most significant SNP (rs17277986) were all significant with the smallest $p$ value $= 0.0005$.

All the interesting SNPs are in intron 1 of SORCS1. This gene is 590 kb, and intron 1 itself is 207 kb and the largest intron in the gene. Most of the regions in intron 1 are conserved (>70% identity) over the 100-bp calculation window among mammals (data not shown). Thus, this region may be important as a functional element in the gene.

Grupe et al. [2006] found an association between SNP rs600879 (164 bp away from rs17277986) in SORCS1 and AD with a
p = 0.0043 in their combined data set. SNP rs600879 (108,913,108 bp) is 608 bp away from the first intron/exon boundary. However, it was not consistently replicated in all of their individual datasets as it was significant in only two of them (p = 0.017 and 0.040). However, the sample size of the homozygotes with the minor allele was small (between four and nine samples for cases and controls) because the minor allele frequency of this SNP was 11.2 and 8.5% for cases and controls, respectively. The SNP was not significant in the other two data sets.

In our study, SNP rs17277986 in SORCS1 was significantly associated with AD in females in the case–control and independent family-based data sets. There are several possible hypotheses for the role of SORCS1 in AD.

First, it could be due to the estrogen effect. Studies on the estrogen-replacement therapy in AD suggested that estrogen may provide some protection against memory loss and lower the risk of developing AD [Burns and Murphy, 1996; Tang et al., 1996; Wickelgren, 2003]. Second, this variant may be in or near intrinsic regulatory sequences that might govern cell type-specific or tissue-specific expression of SORCS1. Third, genomic imprinting may also play a role for the significant difference of the alleles between AD cases and controls, where differential gene expression depends upon whether the inheritance is through the mother or father [Hall, 1990]. Methylation has been proposed as a mechanism of imprinting [Holliday, 1989] and is supported by the findings of the increased number of unmethylated sites in AD patients in comparison to controls [Payao et al., 1998]. Fourth, there may be interaction between mitochondrial DNA mutations and this autosomal locus such that a particular mitochondrial genotype is required for individuals carrying this autosomal risk variant to express disease. It was suggested that mitochondria are involved in apoptosis and there is evidence of oxidative damage to mitochondrial DNA in AD cases [Green and Reed, 1998; Mecocci et al., 1994]. Although there is no evidence that any of these possible explanations are, in fact, correct, they provide plausible areas for further research.

Epistasis, or gene–gene interaction, is crucial in detecting polymorphisms associated with an increased risk of disease [Moore, 2003; Sing et al., 2004; Thornton-Wells et al., 2004]. In the present study, we applied MDR to investigate the potential associations between AD susceptibility and candidate genes covered from linkage, gene expression, and association studies. MDR identified two SNPs (rs1277738 and rs10741083) in CACNB2 as having an interaction effect among genes on chromosome 10. These two SNPs were both intronic SNPs: one was in intron 2 and the other was in intron 5. They are 216 kb apart and not in linkage disequilibrium with each other (r^2 = 0.002). One SNP, rs1277738, had a very strong genotypic association (p = 0.003) with AD and it was identified as having a main effect in MDR analysis. Another SNP, rs10741083, did not show single marker association with AD. However, the interaction effect between this SNP and the main effect SNP had higher prediction accuracy (Table 6), suggesting that the interaction effect was not absolutely driven by rs1277738, which had the main effect.

Table 8. Summary of MDR Results in the Female Subset

<table>
<thead>
<tr>
<th>No. locus</th>
<th>Best model</th>
<th>Accuracy (%)</th>
<th>Crossvalidation</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-locus</td>
<td>SORCS1(rs17277986)</td>
<td>54.74</td>
<td>80%</td>
<td>0.098</td>
</tr>
<tr>
<td>2-locus</td>
<td>SORCS1(rs11193054) CACNB2(rs1277738)</td>
<td>48.67</td>
<td>20%</td>
<td>0.884</td>
</tr>
<tr>
<td>3-locus</td>
<td>SORCS1(R517277986) CACNB2(rs1409207) CACNB2(rs2482100)</td>
<td>52.49</td>
<td>40%</td>
<td>0.297</td>
</tr>
</tbody>
</table>

CACNB2 encodes a voltage-dependent calcium channel beta 2 subunit. The beta 2 subunit works as a complex with other subunit (e.g., alpha unit) and the protein complexes play pivotal roles in signal transduction and homeostasis processes [Opatowsky et al., 2003]. The polymorphisms in the introns may affect the binding site that binds alpha unit. If the interaction between the alpha and beta subunit is affected, the function of the whole protein complex is subsequently affected.

Interpreting results is a challenge in exploring gene–gene interactions. MDR analysis on genes on chromosome 10 had low crossvalidation consistency on the best model (two SNPs in CACNB2), although the interaction was significant in the logistic regression analysis when noise was eliminated from the data set. It may be the case that multiple models are all good predictors of disease risk. If a gene is always in the top MDR models, it also suggests that the gene may have an effect. We discovered three genes (CACNB2, PTPLA, and SORCS1) that showed consistent evidence of association with AD in our previous studies. MDR identified CACNB2 as the single best model, but not PTPLA or SORCS1. By considering the frequencies of specific models in the top 10 MDR models, we were able to find that these three genes were in the top 10 MDR models most of the time. This is another way to identify important effects in the disease pathology. The data suggest that each gene has a modest effect and that an extensive epistasis or gene–gene interaction is underlying the disease etiology.

Our genomic convergence study suggests that genetic variations in PTPLA, SORCS1, and CACNB2 genes may be associated and have modest effect to the risk of AD. In AD, the APP is enzymatically cut to generate Aβ. Aβ accumulates and aggregates to form an oligomer and cause apoptosis, which can cause AD [Hardy and Higgins, 1992]. The SORLA gene is associated with AD and it inhibits the generation of Aβ. SORCS1 is a homolog of SORLA and may have a similar function as SORLA to inhibit the generation of Aβ. CACNB2 is a calcium channel protein and affects the calcium level, which could cause mitochondrial damage and then induce apoptosis [Finlin et al., 2006]. PTPLA is a phosphatase, but relatively little is known about

Figure 4. Hypothesized pathways involved in AD.
its biological function. Base on its phosphatase function, we can conjecture that it is involved in the phosphorylation of the Tau protein; phosphorylated Tau forms neurofibrillary tangles, which is one of the hallmarks of AD. PTPLA might also be involved in the phosphorylation of GSK-3β protein which binds β-catenin. β-Catenin is associated with elevated Aβ42 [Prager et al., 2007; Shim et al., 2007]. β-Catenin can also bind to transcription factors and induce unscheduled cell cycles, which is also hypothesized to be related to AD (Fig. 4). In conclusion, this study suggests that genetic variations in PTPLA, SORCS1, and CACNB2 genes might alter the risk for AD by affecting multiple pathways. The replication of the effect of these genes in different study populations and search for susceptible variants and functional studies of these genes are necessary to get a better understanding of the roles of the genes in AD.

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References


