A Cytogenetic Abnormality and Rare Coding Variants Identify ABCA13 as a Candidate Gene in Schizophrenia, Bipolar Disorder, and Depression

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Schizophrenia and bipolar disorder are leading causes of morbidity across all populations, with heritability estimates of ~80% indicating a substantial genetic component. Population genetics and genome-wide association studies suggest an overlap of genetic risk factors between these illnesses but it is unclear how this genetic component is divided between common gene polymorphisms, rare genomic copy number variants, and rare gene sequence mutations. We report evidence that the lipid transporter gene ABCA13 is a susceptibility factor for both schizophrenia and bipolar disorder. After the initial discovery of its disruption by a chromosome abnormality in a person with schizophrenia, we resequenced ABCA13 exons in 100 cases with schizophrenia and 100 controls. Multiple rare coding variants were identified including one nonsense and nine missense mutations and compound heterozygosity/homozygosity in six cases. Variants were genotyped in additional schizophrenia, bipolar, depression (n > 1600), and control (n > 950) cohorts and the frequency of all rare variants combined was greater than controls in schizophrenia (OR = 1.93, p = 0.0057) and bipolar disorder (OR = 2.71, p = 0.00007). The population attributable risk of these mutations was 2.2% for schizophrenia and 4.0% for bipolar disorder. In a study of 21 families of mutation carriers, we genotyped affected and unaffected relatives and found significant linkage (LOD = 4.3) of rare variants with a phenotype including schizophrenia, bipolar disorder, and major depression. These data identify a candidate gene, highlight the genetic overlap between schizophrenia, bipolar disorder, and depression, and suggest that rare coding variants may contribute significantly to risk of these disorders.

Introduction

The lifetime prevalence of both schizophrenia (MIM #181500) and bipolar disorder (MIM #125480) is about 1% and of major depressive disorder (MIM #608516) around 15%.1,2 With onset in young adulthood and lifelong duration, these illnesses impose a massive burden on individuals, their families, and healthcare providers.3 Although current symptom-based classifications define these as separate mental disorders, the boundaries between diagnoses are indistinct. Thus the incidence of depression and bipolar disorder is increased among relatives of probands with schizophrenia,2,4 and twin studies confirm genetic and environmental correlations between bipolar disorder and major depression.5 Estimated heritabilities of more than 80% for schizophrenia and bipolar disorder6,7 and 40% for recurrent major depression8 have driven the search for causative genes. Various approaches including linkage, gene association studies, investigation of cytogenetic abnormalities, and, recently, the detection of copy number variants (CNVs) have contributed to our understanding of genetic risk in these conditions but have, necessarily, only interrogated particular aspects of the continuum of human genetic variation. Recent genome-wide association studies (GWAS) that probe common variation at the whole genome scale in large case-control cohorts have found, and sometimes replicated, significant associations of a number of candidate genes with schizophrenia and bipolar disorder and have found substantial overlap of risk factors between these two illnesses.9–14 These studies have led to a reevaluation of models of the inheritance of schizophrenia and suggest that common polygenic variation may explain more than one third of the genetic contribution to schizophrenia.11

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However, rare CNVs involving genes also make an important additional contribution to the total genetic variation associated with disease, and a parallel role for rare coding sequence variants seems likely. In particular, the role of rare mutations of large effect (genotype relative risk > 2) remains largely unexplored and is unlikely to be clarified by association strategies via common SNPs.

In the same way that CNVs have proved informative, the study of rare cytogenetic abnormalities can reveal major gene effects on disease risk for the individual and provide useful insights into etiological pathways relevant to illness in the wider population. The genes DISC1, PDE4B, GRIK4, and NAPA3 have each been identified at unique translocation breakpoints in single patients with schizophrenia and subsequently found to be associated with both schizophrenia and bipolar disorder in populations.

In this paper we describe the discovery, through cytogenetics, of a candidate gene, ABCA13, for schizophrenia, bipolar disorder, and depression. ABCA13 is a member of the adenosine triphosphate (ATP)-binding cassette (ABC) super-family encoding transporters shuttling a variety of substrate/allocrites across cellular membranes. Gene mutations in members of the seven ABC subfamilies (A-G) contribute to human disorders with Mendelian or complex inheritance. In higher organisms, ABC proteins typically contain two transmembrane domains (TMDs) and two cytoplasmic nucleotide binding domains (NBDs), the latter containing characteristic ATP-binding Walker A and B motifs. We resequenced ABCA13 gene exons encoding these key functional domains in a “discovery” cohort of 100 cases with schizophrenia and 100 controls and identified 10 nonsynonymous rare coding variants. The frequency of these variants was then determined in a “test” sample of subjects with schizophrenia, bipolar disorder, major depression, and controls. Structural modeling and in silico prediction tools suggested pathological effects. We followed up affected and unaffected relatives of probands with mutations and demonstrated significant linkage of mutations with a phenotype that included schizophrenia, bipolar disorder, and depression. Finally, we determined that ABCA13 protein is expressed in mouse and human hippocampus and cortex, both regions relevant to schizophrenia and bipolar disorder.

Material and Methods

Clinical Description of the Individual Carrying the Complex Chromosomal Rearrangement

The 48-year-old male carrying the complex chromosomal rearrangement inv(7)(p12.3;q21.11), t(7;8)(p12.3;p23) was identified by W.J.M. and was initially described in the hospital case notes as carrying a simple t(7;8) reciprocal translocation. The patient gave full consent to the study and was interviewed on separate occasions by a consultant psychiatrist (W.J.M.) and a psychiatric nurse.

The patient had a diagnosis of severe chronic schizophrenia, with continuous symptoms since his first admission to psychiatric hospital at the age of 16. From his records and discussions with ward staff, there was clear evidence of auditory hallucinations and bizarre delusions of a persecutory and grandiose nature in the past lasting more than 6 months. After an initial series of inpatient stays, he had been continuously in hospital for more than 25 years. On review he showed the classical negative signs of severe chronic schizophrenia with self-neglect to the extent that he needed complete support with self-care, and he showed formal thought disorder and exhibited motor stereotypes with restlessness and frequent pacing. All these symptoms had been constantly present for many years. He received clozapine and haloperidol as treatments for his schizophrenia and procycline to counteract side effects of these. He was on no other medications save a mild laxative. There had been no contact with any members of his family and details of family history were not available.

Clinical Description of Patient and Control Cohorts

The study was approved by the Central Office of Research Ethics Committees (COREC) in Scotland and all subjects gave informed written consent. Subjects were from the Northeast and Southeast of Scotland contacted through in-/out-patient services of hospitals in Scotland, and diagnoses were made according to DSM-IV criteria. All patients were interviewed by an experienced psychiatrist or psychiatric nurse via the “Schedule for Affective Disorders and Schizophrenia-Lifetime” version (SADS-L), supplemented by hospital case note review and information from carers and relatives. Diagnosis was reached by consensus between two psychiatrists.

A family study was designed to: (1) determine whether the rare variants were de novo or familial mutations; (2) describe the range of phenotypes in individuals with variants; and (3) test for linkage between variants and different phenotypes. Attempts were made to contact all first and second degree relatives of the probands who carried one of the ten variants selected for follow up. Some probands were members of large families previously recruited for linkage studies, and when possible these families were revisited so that diagnoses could be confirmed or updated. DNA samples were obtained from 106 relatives of 21 individuals carrying one of the rare mutations. DSM-IV diagnoses of family members were based on direct interview by a psychiatrist as described above.

Controls of similar North European ancestry were recruited through: (1) the South of Scotland Blood Transfusion Service and the Grampian Blood Transfusion service, (2) students taking part in an unrelated study, (3) hospital staff and social networks of the research team, and (4) the 1936 Lothian Birth Cohort (LBC1936). Controls were not screened for personal or family history of psychiatric illness.

These case-control individuals are part of a stable population that shows little evidence of stratification. Moreover, this cohort has been extensively genotyped in case-control studies of candidate genes producing both negative and positive results, suggesting a lack of cohort-dependent frequency bias.

Fluorescence In Situ Hybridization Analysis

BAC and fosmid clones mapping to 8p23.3 and 7p12.3 were selected with the UCSC genome browser database (Table S1 available online). Clone DNA prepared with the QIAGEN miniprep kit was labeled with Biotin dUTP or Digoxigenin dTTP by nick translation and hybridized to patient metaphase spreads with standard FISH methods. Slides counterstained with DAPI in Vectashield antifade solution were assessed on a Zeiss Axioskop2 fluorescence microscope.
microscope with a chroma number 81000 multispectral filter and a Hamamatsu Photonics Orca-ER digital camera with SmartCapture2 software (Digital Scientific, UK).

DNA Resequencing

ABCA13 is a very large gene with 62 exons, so sequencing was restricted to selected exons encoding key functional domains: two NBDs, two TMDs, and a conserved hydrophobic membrane-dipping region (HDR). Oligonucleotide primers were designed with PRIMER3 software against exons 32–35, 37–40, 43, 50–53, and 54–59 of ABCA13 (NM_152701). PhredPhrap and Consed software were used for resequencing analysis. Confirmation of variants was obtained through independent resequencing reactions, restriction enzyme digests, cloning PCR products, and ABI TaqMan assays (all primers listed in Table S2).

Bioinformatics: Splicing, Sequence Alignment, Homology Modeling

Potential splice site variants were assessed with Berkeley Drosophila Genome Project splice site prediction tools.34 Human ABCA13 orthologs were determined with a BLAST search against the translated nonredundant sequence database (tblastn), via the NCBI website with default parameters.32,33 Seven orthologous sequences were retrieved (GenBank identifiers): Homo sapiens (259013576), Pan troglodytes (114613334), Macaca mulatta (109066580), Mus musculus (30023642), Rattus norvegicus (158187532), Canis familiaris (73981662), Bos taurus (194666167), and Gallus gallus (50736039).

Sequences for all NBDs from ABCA transporters were retrieved directly from the SwissProt database.34 Because of their highly conservative sequences, the ortholog and ABCA-NBD multiple sequence alignments were generated automatically with ClustalX55 (Figure S1).

Numerous examples of experimentally determined ABC nucleotide-binding domains are deposited in the Protein Data Bank (PDB).36 For modeling human ABCA13-NBD1 (SwissProt Accession No: Q86UQ4; amino acid residues 3842–4074), three of its closest experimentally determined homologs, ascertained by a BLAST search against the PDB, were used as templates. These included X-ray structures from Thermatoga maritima (PDB ID: 1VPL; 2.1 Å) (Joint Center for Structural Genomics, unpublished), CysA from Alcyoncidibacillus acidocaldarius (PDB ID: 1247, chain A; 1.9 Å), and MalK from Escherichia coli (PDB IDs: 1Q12, chain A).37 The higher resolution 1Q12 (2.6 Å) structure of MalK E. coli was selected instead of 1Q1E (2.9 Å) and 1Q1B (2.8 Å). The target ABCA13-NBD1 shares 31%, 29%, and 30% sequence identity, respectively (~52%–54% similarity), when compared with the templates from N-to-C terminus.

The alignment between the target and template sequences for modeling purposes was based on an initial multiple sequence alignment among them and a range of other-related protein sequences with the program MUMMALS38 (Figures S2 and S3).

Related sequences were first identified via the Network Protein Sequence Analysis server, with a BLASTSearch against the SwissProt database.34,39 After removing identical sequences from the returned output and considering only those sequences with at least 200 amino acids in length, 413 protein sequences were identified, and their corresponding homologous NBD-like protein regions were extracted (E-value cut-off ≤ 1e-10) and aligned with MUMMALS (Hiddden Markov model option: HMM_1.3.1; Identity threshold: 0.6). The target-template alignment was further manually refined from the multiple sequence alignment in order to place gaps optimally guided by positioning of secondary structure elements, identified by STRIDE for the templates, and predicted by PsiPred version 2.5 for the target sequence.40–42

The program Modeler release 8 version 2 was used for model building based upon the aforementioned alignment.41 Twenty models were generated, and the one with the lowest objective function score was selected as the representative model. Nonidentical side chain residues (between target and templates) for the representative model were optimized with the side chain replacement program, SCWRL version 3.44,45 The model was then protonated under SYBYL version 6.9 (Tripos Associates, St. Louis, MO) and subject to brief energy minimization (30 steps steepest descent followed by 30 steps conjugate gradients) employing the Tripos forcefield to remove clashes and bad geometries.46 The model was finally checked for valid stereochemistry with PROCHECK version 3.5.4 (most favored regions, 89.8%; additional allowed regions, 9.8%; generously allowed regions, 0%; disallowed regions, 0.5%) and validated with the MetaMQAP II server47 (GDT_TS, 69; RMSD, 3.3 Å).47 All surface area calculations were performed with GETAREA version 1.1 via the following parameters: radius of water probe in Angstroms = 1.4; output level = Area per residue.48 The hydrogen-bonding network was analyzed with SYBYL version 6.9 (Tripos Associates) and the Protein Interactions Calculator.49

Statistical Analysis

Variants for further study were selected from resequencing “discovery” set data. However, all statistics relating to individual and global variant influence were based solely on TaqMan genotyping data derived from the secondary “test stage.” This approach ensured unbiased estimates of effect sizes. For each variant in the test stage, allele frequencies were compared with Fisher’s exact test. A combined association test across all nonsynonymous mutations was performed by comparing the number of variants in cases to the number in controls. To account for variable sample size across the total, sample size was adjusted to N = n / (Σ(1/Ni)), where Ni is the samples size at the ith variant and n is the number of variants. The number of observed variants was adjusted as Σ(pi)N, where pi is the frequency of the ith variant. The population attributable risk (PAR) was calculated as follows: PAR(%) = 100 * (P(any ns variant) * (OR – 1) / [1 + P(any ns variant) * (OR – 1)] with P(any ns variant) = the probability of having any of the ns variants, calculated from the controls. This probability (~ 0.024) is the same for all groups because it is calculated from the controls.

Nonparametric linkage analysis was conducted with Merlin50 to calculate allele sharing between all sets of affected relatives in the 14 families informative for linkage with two or more affected relatives, under both a “narrow” phenotype definition (schizophrenia and bipolar disorder) and a “broad” phenotype (schizophrenia, bipolar disorder, and major depression). For each SNP, the allele frequency on which identity-by-descent is estimated was specified to be 0.0025 (average mutation rate in controls). The LOD score and p value were calculated via the exponential model described by Kong and Cox.51

Immunofluorescence

Two separate Abca13 antibodies, NC1 and 807, were raised in rabbits via conjugated peptide immunization. Frozen adult mouse
brain sections (10 μm thickness) were fixed in ice-cold acetone and blocked, incubated with primary antibody, and washed in phosphate-buffered saline + 0.1% Tween. After Alexa Fluor fluorescent donkey anti-rabbit secondary antibodies incubation and washing, slides were mounted in DAPI/antifade solution. Images were captured as described for FISH.

Results

Cytogenetics

We characterized a chromosomal rearrangement, inv(7)(p12.3;q21.11), t(7;8)(p12.3;p23), comprising a pericentric inversion of chromosome 7 coupled with a translocation between chromosomes 7 and 8 in a patient with chronic schizophrenia. FISH confirmed only one break-point, at 7p12.3, directly disrupted a gene, ABCA13 (Figures 1A and 1B; Table S1). We hypothesize that ABCA13 disruption causes illness in this individual through haploinsufficiency that commonly occurs in similarly truncated transcripts through nonsense-mediated decay.

Resequencing Functional Domain Exons

The ABCA13 locus had not previously been associated with psychiatric disorders. To investigate whether rare functional variants in this gene contribute to psychiatric disorders, we resequenced 19 out of the total of 62 exons in the discovery cohort of 100 unrelated cases with schizophrenia (the same diagnosis as the patient with the chromosome abnormality) and 100 controls. These exons were selected to encode key functional domains: two NBDs, two TMDs, and a conserved hydrophobic membrane-dipping region (HDR) (Figure 1C).

We identified 32 variants not previously recorded in dbSNP: 20 in exons and 12 within flanking intronic sequences (Table 1; Figure S4). Of the 20 exonic variants, 14 were nonsynonymous, of which 8 were identified initially in schizophrenia cases, 3 in schizophrenia and controls, and 3 in controls only.

The Frequency of Variants in Cases and Controls

Ten variants were selected for further analysis in a test cohort of subjects with schizophrenia (1019), bipolar disorder (680), major depression (365), and 2270 controls, based on the criteria: (1) present in no more than one of 100 controls, (2) causing a nonconservative amino acid change, and (3) showing cross-species conservation of the wild-type residue (Table 2). Genotyping assays designed against these 10 mutations were used to screen cases and controls. Despite relatively small numbers, three variants (H3609P, T4031A, and T4550A) were significantly more frequent in bipolar than controls and R4843L was more common in schizophrenia (Table 2), although none remained significant after correction for multiple testing of 10 variants in three phenotypes. A global comparison of the frequencies of all ten variants between cases and controls in this test stage showed a significant increase in the frequency of mutations in schizophrenia (p = 0.0057 OR = 1.93 95% CI 1.24) and bipolar disorder (p = 0.00007 OR 2.71 95% CI 1.73) but only a trend in major depression (p = 0.096 OR 1.67 95% CI 0.88). Similar significance was obtained when all case diagnoses were grouped (p = 0.0004 OR 2.1 95% CI 1.45). The population attributable risks of the variants were 2.21%, 4.00%, and 2.61%
for schizophrenia, bipolar disorder, and across all diagnoses, respectively.

**Study in Families of Individuals with Mutations**

Relatives of probands carrying eight of the ten rare mutations gave consent to take part in the family study designed to investigate the phenotypes associated with mutations and to determine whether these were de novo or familial mutations. 106 relatives in 21 families were genotyped (Figure 2). Families ranged in size from 2 to 16 persons and the diagnoses in family members (including probands) carrying one of the variants were schizophrenia (14), bipolar disorder (10), major depression (19), minor depression, anxiety, or alcoholism (3), and no psychiatric diagnosis (19). No variants were confirmed as de novo mutations.
Table 2. The Properties of Ten Rare ABCA13 Variants Genotyped in Large Test Cohorts of Cases and Controls

<table>
<thead>
<tr>
<th>Variant Domain</th>
<th>Exon</th>
<th>Genomic Position</th>
<th>Base Change</th>
<th>WT Residue Conservation</th>
<th>Residue Change</th>
<th>Type</th>
<th>Representation in Discovery Set</th>
<th>SCZ (n = 100)</th>
<th>Controls (n = 100)</th>
<th>SCZ by Diagnosis (Total)</th>
<th>Control (Total)</th>
<th>SCZ p Value</th>
<th>SCZ Odds Ratio</th>
<th>BP p Value</th>
<th>BP Odds Ratio</th>
<th>MDD p Value</th>
<th>MDD Odds Ratio</th>
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<tbody>
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<td>TMD1</td>
<td>33</td>
<td>48,382,318</td>
<td>G→A</td>
<td>RH, D, R, M</td>
<td>R3604Q</td>
<td>missense</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0/841 SCZ, 0/494 BP, 0/331 MDD</td>
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<td>(951)</td>
<td>0.334</td>
<td>1.37</td>
<td>0.050</td>
<td>2.43</td>
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<td>RH, D, R, M, Ch</td>
<td>H3609P</td>
<td>missense</td>
<td>3</td>
<td>1</td>
<td>25</td>
<td>10/857 SCZ, 10/488 BP, 5/262 MDD</td>
<td>8</td>
<td>(939)</td>
<td>0.334</td>
<td>1.37</td>
<td>0.050</td>
<td>2.43</td>
<td>0.130</td>
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<td>48,382,619</td>
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<td>S3704R</td>
<td>miss./splice</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0/851 SCZ, 0/496 BP, 0/259 MDD</td>
<td>0</td>
<td>(939)</td>
<td>0.334</td>
<td>1.37</td>
<td>0.050</td>
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<td>0.130</td>
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<td>RH, D, R, M, Ch</td>
<td>T4031A</td>
<td>missense</td>
<td>1</td>
<td>0</td>
<td>3</td>
<td>0/1019 SCZ, 3/678 BP, 0/365 MDD</td>
<td>0</td>
<td>(2262)</td>
<td>0.334</td>
<td>1.37</td>
<td>0.050</td>
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<td>H4262R</td>
<td>missense</td>
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<td>0</td>
<td>0</td>
<td>0/837 SCZ, 0/562 BP, 0/318 MDD</td>
<td>0</td>
<td>(980)</td>
<td>0.334</td>
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<td>0.050</td>
<td>2.43</td>
<td>0.130</td>
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<td>T4550A</td>
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<td>1</td>
<td>0</td>
<td>18</td>
<td>9/861 SCZ, 8/482 BP, 1/265 MDD</td>
<td>3</td>
<td>(938)</td>
<td>0.334</td>
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<td>0</td>
<td>1</td>
<td>17</td>
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<td>(971)</td>
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<td>1.5</td>
<td>0.087</td>
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<td>P4648A</td>
<td>missense</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>0/869 SCZ, 0/619 BP</td>
<td>0</td>
<td>(959)</td>
<td>0.38</td>
<td>1.5</td>
<td>0.087</td>
<td>2.75</td>
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<td>48,534,520</td>
<td>C→T</td>
<td>RH, D, R, M</td>
<td>R4728X</td>
<td>nonsense</td>
<td>1</td>
<td>0</td>
<td>10</td>
<td>4/1004 SCZ, 5/680 BP, 1/374 MDD</td>
<td>5</td>
<td>(2270)</td>
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<td>1.51</td>
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<td>48,597,317</td>
<td>C→T</td>
<td>RH, D, R, M</td>
<td>R4843C</td>
<td>missense</td>
<td>0</td>
<td>1</td>
<td>19</td>
<td>12/815 SCZ, 5/619 BP, 2/336 MDD</td>
<td>6</td>
<td>(1025)</td>
<td>0.047</td>
<td>2.54</td>
<td>0.400</td>
<td>1.38</td>
<td>0.630</td>
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</table>

Global significance of all variants by diagnosis (95% CI lower limit shown for odds ratios):

- SCZ: 5.70E-03 1.93 (1.24)
- BP: 7.42E-05 2.71 (1.73)
- MDD: 9.66E-02 1.67 (0.88)

Population attributable risk by diagnosis:

- SCZ: 2.21%
- BP: 4.00%
- MDD: 0.02%
Nonparametric single point linkage analysis with MERLIN\textsuperscript{50} to estimate identity by descent (IBD) between all pairs of affected relatives in the 14 multiply affected families informative for linkage gave a combined Z score of 4.18 and a Kong and Cox\textsuperscript{51} LOD score of 4.38 ($p = 3.5 \times 10^{-6}$), establishing significant linkage of $\text{ABCA13}$ mutations with a “broad” phenotype that included cases with schizophrenia, bipolar disorder, and major depression. A LOD score of 2.1 ($p = 0.0009$) was achieved when the analysis was performed under the more restrictive phenotype definition including only cases with schizophrenia and bipolar disorder.

Genetic Properties and Clinical Phenotypes Associated with Individual Rare Variants

Data on each variant are presented in Table 2. Figure 3 highlights the sequence conservation and structural consequences of the potential null mutations R4728X and S3704R and missense mutation T4031A. Figure S5 shows the location of the nine predicted pathological variants on a topological representation of ABCA13 protein.

1. R3604Q was detected in a single discovery case with schizophrenia and one of 951 controls. A pathological effect was predicted by pmut and Panther online utilities (Table S3).\textsuperscript{52–55} In the family of the schizophrenia proband (Fam19), the mutation was present in a parent and two siblings and all three relatives had affective disorder.

2. H3609P was detected in three subjects with schizophrenia and one control in the discovery cohort and was more frequent in bipolars than controls ($p = 0.05$ OR 2.43) in the test cohort. A pathological effect was predicted by pmut. Eight families were followed up and the mutation was detected in four cases of bipolar illness or depression in Fam17, three siblings with schizophrenia in Fam11, two cases of schizophrenia, bipolar disorder, or depressive episodes in each of families 12, 14, and 16, and in an unaffected parent in families 13, 15, and 18. Three bipolar cases were compound heterozygotes carrying both the T4550A and H3609P mutations and one person with severe chronic schizophrenia was homozygous for H3609P whereas all controls carrying variants were monoallelic, reinforcing the possibility of interactive effects of rare variants on these phenotypes.

3. S3704R, located in an extracellular loop of the first TMD (Figures 3C and 3F), was identified in a subject with schizophrenia in the discovery cohort and cosegregated with schizophrenia (one case) and depression (four cases) in the proband’s family (Fam10). The mutation was not detected in a further 1606 cases or in 939 controls. A pathological effect was predicted by pmut. In addition to its amino acid substitution, this mutation introduces a novel cryptic splice donor site, potentially resulting in a transcript with a downstream frameshift and premature stop codon. Hence, S3704R displays the features of a rare Mendelian, highly penetrant, deleterious mutation.

4. T4031A, a missense variant located in NBD1 (Figures 3B and 3E), was identified in three individuals with bipolar

<table>
<thead>
<tr>
<th>P Value</th>
<th>Odds Ratio</th>
<th>Odds Ratio</th>
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<tbody>
<tr>
<td>Global significance of all variants across all diagnoses (95% CI lower limit shown for odds ratio):</td>
<td>2.60E-04</td>
<td>2.1 (1.45)</td>
</tr>
<tr>
<td>Population attributable risk across all diagnoses:</td>
<td>0.0%</td>
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The genomic location, nature of the nucleotide mutation, resulting amino acid change, and cross-species conservation of the wild-type residue are listed. The statistical significance (significant p-values highlighted in bold type), odds ratio, and population attributable risk for individual and grouped (global assessment) variants are detailed in relation to their frequency in the individual and combined case and control groupings. In conjunction with pathological prediction and family data, only P4648A failed to show evidence for a causal role in psychiatric illness. Genetic position based on NCBI build 36. p-values calculated as follows: pmut = 1 - (p-value for dominant disorder) $\times$ (p-value for recessive disorder).
disorder in a sample of 2072 cases and was absent in 2262 controls \( (p = 0.012) \). A pathological effect was predicted by Panther. As with R4728X (see below), the missense variant occurred on a distinct haplotype \( \text{(Table S4)} \), implying that present-day carriers of the mutation are likely to be descended from a common European ancestor.

T4031A is strictly conserved across species and among all human ABCA NBD1s \( \text{(Figure 3E; Figure S1)} \), and we evaluated the consequence of the mutation on protein structure and function by means of a comparative three-dimensional model \( \text{(Material and Methods; Figures 3G and 3H)} \). T4031 lies within the protein core immediately adjacent to the functional H-loop histidine residue \( (H4032) \) and in proximity to other functional motif residues from Walker A \( (G3875) \) and Walker B \( (E4001) \). Substitution to Ala likely alters H-loop conformation and, in turn, catalytic activity. The adjacent H-loop histidine residue \( (H4032) \) is essential for ATP hydrolysis in NBDs \( ^{56,57} \) and the equivalent mutation, H2128R, in NBD2 of human \( \text{ABCA4} \) has been implicated in Stargardt disease \( (\text{MIM} \ #248200) \), a lipid deposition disorder.\( ^{58} \)

5. H4262R, located distal to the HDR, was detected in a single person with schizophrenia and an unaffected sibling \( \text{(Fam22)} \) but was absent from control, bipolar disorder, and depression cohorts. A pathological effect was predicted by pmut and PolyPhen.

Figure 2. Cosegregation of ABCA13 Variants with Illness in Families
These families were all included in the linkage analysis. Images created with CraneFoot application.\(^{83}\)
6. T4550A is a missense variant found more frequently in bipolars than controls (p = 0.0097 OR 5.52) with a similar trend in schizophrenia (p = 0.054 OR 3.29) but not depression (p = 0.63 OR 1.18). Two families were followed up and the mutation was detected in a second sibling with schizophrenia in one family (Fam20) and an unaffected sibling in the other (Fam21).

7. R4590W was discovered in a control and in the test stage was relatively frequent and found in 17 cases and 4 controls. A pathological effect was predicted by pmut and PolyPhen. No families were available for follow-up.

8. P4648A was discovered in a single control and was not detected in the test samples. Controls were not screened for current or past episodes of mental illness and family details are not available. We consider this variant non-pathological.

9. R4728X, a nonsense mutation located at the start of NBD2 (Figures 3A and 3D), was found in four cases with schizophrenia, five with bipolar disorder, one with depression, and five controls (not screened for personal or family history of psychiatric illness). Four of the mutation carriers had relatives who agreed to take part in the study. In two of these families (Fam3 and Fam4), the mutation segregated with bipolar disorder and depression and in two (Fam5 and Fam6) with schizophrenia and depression (Figure 2). The mutation showed incomplete penetrance and was found on a distinct haplotype background (as T4031A, see above) (Table S4). The functional significance of the mutation is not known but transcripts containing nonsense mutations are commonly degraded, resulting in reduced gene dosage. Moreover, an equivalent mutation, R1275X, within the conserved A-loop motif in the human ABCC6 gene is pathological, causing pseudoxanthoma elasticum (MIM #264800).

10. R4843C was initially discovered in a control, but in the test stage was more frequent in cases. A pathological effect was predicted by pmut. The comparison of schizophrenia (12/815) with controls (6/1025) was significant (p = 0.047 OR 2.54). Follow-up was possible with three families where the mutation was present in a bipolar parent in Fam7, an unaffected parent in Fam8, and a sibling pair with depression Fam9. A bipolar parent of the proband with unipolar depression in Fam9 did not have the mutation but we have no diagnosis for the other parent who has the variant.

Compound Heterozygosity

Strikingly, five cases were found to be compound heterozygotes and one homozygous, whereas all controls carrying variants were monoallelic, suggesting possible additive effects on risk. Two of the four cases with T4031A, one with schizophrenia and one bipolar, were compound
heterozygotes also carrying the variant T4550A. Only one relative of the two compound heterozygotes was available for study and this unaffected parent of the proband with schizophrenia carried only T4031A. Two bipolar cases were compound heterozygotes carrying both the H3609P and R4590W mutations, one bipolar case carried H3609P and T4550A, and one person with severe chronic schizophrenia was homozygous for H3609P.

**Abca13 Expression in Human and Mouse Brain**

RT-PCR confirmed earlier reports of *ABCA13* mRNA transcript expression in brain and indicated specific expression in human hippocampus and cortex (data not shown). Immunological detection of Abca13 protein by two independently raised antibodies showed low-to-moderate expression in most adult mouse brain regions and higher expression in dentate gyrus granule cells and ventricular areas including choroid plexus and ependymal cells (Figure 4).

**Discussion**

We have identified a cytogenetic disruption of the *ABCA13* gene and 10 rare nonsynonymous mutations of which 9 demonstrated evidence consistent with a role in the etiology of schizophrenia and bipolar disorder. Association of *ABCA13* coding variants with schizophrenia and bipolar disorder was significant for all variants taken together and for specific individual variants. The population attributable risk of the 10 rare variants detected was 2.2% for schizophrenia and 4.0% for bipolar, suggesting that this gene may have an important role in a subgroup of patients and an influence that crosses traditional diagnostic categories. To our knowledge, this is the first instance where multiple and combined coding variants in a single gene have been associated with risk of schizophrenia and bipolar disorder.

*ABCA13* had not previously been implicated as a risk factor in major psychiatric disorders although a suggestive role for common *ABCA13* gene variants in schizophrenia recently emerged from the International Schizophrenia Consortium GWAS data set where a significance level of \( p < 0.001 \) was reached at a single SNP within *ABCA13* and a cluster of SNPs within a few kb of the gene.\(^10\)

The family study included affected and unaffected relatives from 21 families, and in all cases the variants were familial with no instances of de novo mutation. Association with major depressive disorder was not significant in the case-control study but this was a small and conservative group because controls were not screened for personal or family history of psychiatric illness and the control groups would be expected to include some individuals with depression that has a lifetime prevalence of ~15%.
However, a striking finding was that the same mutation was associated with schizophrenia, bipolar disorder, as well as major depression in separate families. The nonsense mutation R4728X was detected in cases with bipolar disorder and depression in Fam3 and Fam4 and with schizophrenia and major depression in Fam5 and Fam6. A similar picture was found for H3609P, which was present in three siblings with schizophrenia in Fam11 and two cases with bipolar disorder and two with major depression in Fam17. The family data also demonstrated incomplete penetrance for seven of the eight variants because these were detected in unaffected relatives. This variable penetrance and range of phenotypes in families with rare ABCA13 mutations is similar to the pattern of illnesses in relatives carrying a translocation disrupting the gene DISC1 who were diagnosed with schizophrenia, depression, and bipolar disorder.61

Rare variants in other members of the ABCA subfamily also give rise to variable disease phenotypes, for example ABCA162 (two lipid-related conditions: Tangier disease [MIM #205400] and familial hypoalphalipoproteinaemia [MIM #604091]) and ABCA463 (four forms of degenerative retinal dystrophy: Stargardt disease, retinitis pigmentosa [MIM #601718], cone rod dystrophy [MIM #604116], and age-related macular degeneration [MIM #153800]), and it has been suggested that precise clinical presentation may be determined by the site of the mutation.62,64 However, in the present study both missense and null mutations contribute to a full range of phenotypes with no clear effect of the site or class of mutation on clinical outcome. The discovery of five compound heterozygotes and one homozygous mutation carrier suggests that additive and interactive combinations of mutations may contribute to these complex phenotypes.

All ABCA proteins functionally characterized to date shuttle lipid molecules across cell membranes.65 More than 1000 different lipid species exist in eukaryotic cells and in addition to being structural components of cellular membranes, different species have key roles in vesicular trafficking, signal transduction, and transcriptional regulation.66,67 Although the precise allocrite for the ABCA13 transporter has not been identified, the presence of a conserved C-terminal motif known to be shared among the lipid transporting group suggests that it is likely to be lipid related.68 Abnormalities in lipid metabolism have been strongly implicated in psychiatric illness with the most consistent finding being depletion of essential fatty acids in tissues of schizophrenic and bipolar patients.69–74 The oligodendrocyte lineage transcription factor 2 (OLIG2) showed association with schizophrenia,75,76 and a growing body of evidence based on imaging studies, microarray analysis, and metabolic and transcriptome investigation has implicated oligodendrogial pathophysiology and myelin biosynthesis in psychiatric disorders.77–79 Consistent with these discoveries are the findings that membrane phospholipids and the expression of genes encoding enzymes important in sphingolipid metabolism are signifi-

antly altered in postmortem brain tissue from schizophrenic patients and upregulated in response to antidepressant drugs.80,81 Disruption of lipid membrane structures and lipid biosynthetic pathways has the potential to affect many important biological processes including intracellular signaling pathways and neurotransmitter function.

Rare sequence variants are known to have an important role in the inheritance of common diseases, and different categories of “rare variant” have been described. Classical rare deleterious familial mutations typically show strong familial aggregation, have high penetrance, and low frequencies. A second type of variant (mostly missense mutations), have higher frequencies of up to 2%–3% in the case population, ORs of 2 or more, and tend to show less familial aggregation.82 In the present study S3704R conforms to the first class of null private mutation with high penetrance and low frequency whereas the missense mutations H3609P, T4550A, and R4590W show characteristics more typical of the second class by virtue of also being detected in controls, having moderate ORs, and higher observed frequencies. These data lend support to a genetic architecture of schizophrenia, bipolar disorder, and depression comparable to other common genetically complex and heterogeneous diseases and provide a novel biological pathway for investigation.

Supplemental Data

Supplemental Data include five figures and four tables and can be found with this article online at http://www.cell.com/AJHG.

Acknowledgments

We would like to dedicate this paper to Walter Muir whose sudden death happened soon after this paper was written. His astute clinical observation and cytogenetic skills were the starting point for studying ABCA13 and his deep understanding of patients with psychosis combined with a formidable knowledge of psychiatric genetics made him an inspiring colleague, mentor, and friend.

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Web Resources

The URLs for data presented herein are as follows:


