Meta-analysis of genome-wide association data detects a risk locus for major mood disorders on chromosome 3p21.1

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Abstract

The major mood disorders, which include bipolar disorder (BD) and major depressive disorder (MDD), are substantially heritable, but few risk loci have been identified. We performed a meta-analysis of 5 major mood disorder case-control samples, including over 13,600 unique individuals genotyped with approximately 500,000 to 1 million single nucleotide polymorphism (SNP) markers on high-density arrays. Allele-wise association results were meta-analyzed with a method...
that weights results by sample size. We found genome-wide significant evidence that SNPs in a region of chromosome 3p21.1 were associated with major mood disorders. The SNP rs2251219 returned the smallest meta-analysis p-value, $3.63 \times 10^{-8}$, with a pooled odds ratio of 0.87. Supportive results were observed in 2 out of 3 independent samples tested in a replication study. These results implicate one or more genes in this region in the etiology of major mood disorders and suggest that BD and MDD share genetic risk factors.

**Keywords**

bipolar; depression; GAIN; STAB1; NT5DC2; PBRM1; NEK4; SPCS1; GNL3; GLT8D1; ITIH

The major mood disorders have a total lifetime prevalence up to 20%, and may soon become the leading cause of morbidity worldwide.1 Similarities in symptoms and treatment response, twin concordance, and shared familial risk (reviewed in ref. 2), have long fed the suspicion that the major mood disorders share genetic risk factors, but molecular evidence remains scarce. Genome-wide association study (GWAS) data offer the opportunity to take a fresh look at genetic factors involved in these common disorders.

Samples (see Table 1) consisted of cases with a major mood disorder and controls, all of European ancestry, ascertained and genotyped as previously described (see Methods). One published GWAS of MDD 3 was not included in the meta-analysis, because complete results were not available, but key SNPs were tested later in a replication study. SNP genotype data were obtained from dbGaP (National Institute of Mental Health BD [NIMH-BP] and Genetic Association Information Network [GAIN-MDD] samples), the Wellcome Trust Case Control Consortium (WTCCC) sample, and through collaborators (German sample). Data from the Systematic Treatment Enhancement Program for Bipolar Disorder (STEP-BD) sample were obtained from the authors’ website. Data from the NIMH-BP, GAIN-MDD, and German samples were used to impute genotypes for about two million HapMap Phase 2 markers. The WTCCC and STEP-BD samples were both genotyped on the same platform, so only the observed data were used.

A total of 317,889 markers could be reliably scored across 4 or 5 samples (Figure 1a). The final results revealed no evidence of residual bias, with a mean genome-wide Z-score of 0.005, close to the theoretical null value of 0 (Supplementary Fig 1). Complete meta-analysis results are presented in Supplementary Table 1.

Six SNPs on chromosome 3p21 were associated with major mood disorder at the $p < 7.2 \times 10^{-8}$ level (Figure 1b; Supplementary Table 2). This corresponds to a genome-wide corrected $p<0.05$ in samples of European ancestry.4 At the most significant marker, rs2251219, the C-allele was consistently less common in cases than controls ($p$-value = $1.12 \times 10^{-8}$; Table 2).

Observed genotypes at rs2251219 were available in all samples except for GAIN-MDD. For that sample, we used imputed genotypes at rs2251219, but the imputation is expected to be highly reliable. Imputation methods perform well in European-ancestry samples, and there was strong linkage disequilibrium (LD) among the markers flanking rs2251219 ($r^2 \geq 0.98$). One nearby SNP that was genotyped in the GAIN-MDD sample, rs2289247, returned a meta-analysis $p$-value of $8.96 \times 10^{-7}$, similar to rs2251219. Random masking of 5% of genotypes had little effect on the imputation results (data not shown). Imputation statistics are presented in Supplementary Table 2.

The analysis of rs2251219 was repeated under a random effects model, with similar results ($p=3.62 \times 10^{-8}$; Supplementary Table 3). The pooled OR was 0.87 (95% CI: 0.83–0.92),
with no evidence of heterogeneity (Q=2.93, df=3, p=ns; I² = 0). GAIN-MDD returned a similar OR of 0.90 (95% CI: 0.82–0.99), consistent with a similar contribution of this locus to both BP and MDD.

To assess robustness, we repeated the analysis 5 times, removing one sample each time. Random effects p-values at rs2251219 ranged from 10^{-6} – 10^{-9}, reflecting varying sample sizes, but the ORs remained stable. (Supplementary Table 3). Thus it appears that the results are not driven by any single sample.

We subsequently obtained association results at rs2251219 in an independent BP case-control sample (n=1536) genotyped by GlaxoSmithKline (GSK). The C-allele of rs2251219 was significantly under-represented in BP cases compared to controls (p=0.002, OR = 0.57; Table 2), replicating our main meta-analysis result. Data from two independent MDD case-control samples were also obtained from GSK: A clinical sample from Munich (n=1792) and a population-based sample from Lausanne (n=1349). No significant association with rs2251219 was found in either MDD sample, although in the larger one the 95% CI of the OR (0.83–1.77) substantially overlapped with that of the present study (0.83–0.92, Table 2). Additional, nearby markers also showed evidence of association in one or more samples (data not shown). When all 3 samples were combined with our original results, the evidence of association at rs2251219 increased (fixed effects p = 1.67 × 10^{-9}; random effects p=4.99 × 10^{-9}). These data provide support for our findings in independent samples, but the findings are more robust in BP than in MDD.

The association signals on chromosome 3p span a ≥246 kb region containing several annotated transcripts (Figure 1b). SNP rs2251219 is a synonymous variant in the gene PBRM1, which encodes polybromo-1, important in chromatin remodeling. The nearby SNP rs2289247 is a non-synonymous (V → M) variant in the gene GNL3, encoding the GTP-ase nucleostemin, involved in proliferation of stem cells, especially in the central nervous system. These genes are good biological candidates, but the LD across the locus is very strong (r² > 0.9; Figure 1c), complicating efforts to localize functional marker(s) in individual genes by association mapping alone.

To help prioritize genes for further study, we examined gene expression in brain tissue. PBRM1 was over-expressed in the dorsolateral prefrontal cortex of patients with BP (p=0.018; Supplementary Figure 2), compared to healthy controls. This finding, not confounded by linkage disequilibrium in the region, independently implicates PBRM1, at least in BP. We also mined 2 brain expression datasets to test 22 disease-associated SNPs for cis-association with expression of all known transcripts in the 3p region. Apparent associations between rs2251219 and expression of PBRM1 and GLT8D1 involved probes that overlap common SNPs, an important source of artifact. Neither were confirmed by qPCR in the Stanley Brain Foundation Samples, and the GLT8D1 finding could not be confirmed in cDNA from the original study using probes that did not overlap known SNPs (data not shown).

This is the first psychiatric study to find genome-wide significant association on chromosome 3p. Previous studies have detected suggestive evidence of association in this region. A meta-analysis of BP GWAS in samples that overlap with those in the present study found suggestive signals; the most significant SNP in that study was nominally significant in the present study (rs1042779; p=0.001). Considering the strong local LD, all of these studies may have detected signals arising from the same risk allele(s).

Previous major mood disorder GWAS have highlighted several genes elsewhere in the genome. We detected association with most of these genes (Supplementary Table 4), but our results are not independent replications since the samples we used overlap with those in the...
previous studies. No signal was significant after genome-wide correction for multiple testing. If many genes play a role in risk, signals may vary from study to study, reflecting small sample differences and other unmeasured factors. Thus, even this large sample may be too small or heterogeneous to detect all important risk alleles at genome-wide significance. A complementary analysis is expected in the future, when the Psychiatric GWAS Consortium13 completes its study of fully-imputed data in these and other samples.

We report molecular support for the prior epidemiologic evidence of genetic overlap between BP and MDD2. These data do not explain why, among all those carrying risk allele(s), some develop BP, others develop MDD, and still others remain apparently well. This phenomenon may reflect a large number of risk alleles, few of which have been detected to date, environmental influences and -- perhaps -- epigenetic factors.

The genetic association findings to date seem to account for little of the inherited risk for mood disorders. Since GWA studies usually omit SNPs with minor allele frequencies below 3–5%, we can say nothing about alleles in that frequency range, even if they have relatively strong effects. If rare alleles of large effect exist, each can account for only a small proportion of cases. More common autosomal alleles (frequency >20%) conferring a heterozygote relative risk of 1.3 would have been detected with >90% power in this sample. Since we found no such alleles, they probably do not exist in this sample, the largest mood disorder sample studied to date. Many loci may add together to confer risk, as some studies suggest14-15, or fewer loci may interact, but to our knowledge, strong epistasis has not yet been demonstrated in complex human traits.

The genetic architecture of the major mood disorders appears to be multi-genic and/or highly heterogeneous. As robust findings accumulate and sample sizes grow, the identified genes may triangulate pathways of etiologic relevance. The GWAS remains an important means to that end, even though each individual finding may represent only a small step forward.

METHODS

Study Samples

The samples used in the meta-analysis have been described previously16–19. Details, including quality control procedures applied to the genotyping data and a description of the replication samples, is provided as a supplement to the published report (Supplementary Note).

Imputation

Genotype data from the NIMH-BP, GAIN-MDD, and German samples were used to impute data on 2.1 million HapMap Phase 2 SNPs by use of the program Markov Chain Haplotyping (MACH), version 1.0 20. MACH uses Markov chain haplotyping to resolve haplotypes, and thereby missing genotypes, from observed genotypes in unrelated individuals. We used the “greedy” algorithm, as recommended by the authors. SNPs that were flagged as having different alleles than in HapMap CEU or as monomorphic were reviewed, after which they were either recoded for the reverse strand (flipped) or dropped. SNPs that were flagged for allele frequencies that were markedly different from HapMap CEU were also reviewed. Palindromic SNPs whose allele frequencies were consistent with reversed coding were flipped. Other SNPs with unexpected allele frequencies were dropped. PLINK21 (vers. 1.4) was used to flip and drop SNPs as necessary. After all allele-coding, monomorphism, and palindrome issues were resolved, imputation was run again. SNPs in the results files were dropped if the MAF in cases or controls was <0.05 or if the error rate...
(as reported in the .erate output file) was >0.01. Finally, the imputed data were formatted into PLINK binaries for analysis.

Meta-analysis

PLINK output (.assoc) files were modified with columns for direction of association, sample size, and strand. For most samples, sample size equaled the sum of cases and controls included in the final analysis, after quality control. For the STEP-BD sample, sample size was set to equal the number of cases only. This was done to avoid over-weighting the results from the NIMH control sample, largely overlapping portions of which were included in both the NIMH bipolar disorder and STEP-BD samples. Modified files were loaded into Metal (July, 2008 version), then processed using the GENOMICCONTROL option, which applies a genomic control correction in samples where the genomic inflation factor is greater than 1.0. Metal weights each sample based on the square root of the sample size.

Great care was taken in combining results from different samples and platforms to avoid confusing alleles, especially at palindromic SNPs. To check this, we inspected all SNPs whose range of allele frequencies in the meta-analysis was greater than 0.2, a commonly-used threshold. Most of these SNPs were palindromes and, as expected, their minimum and maximum allele frequencies across the study samples added to approximately 1. Using the “STRANDLABEL” and “USESTRAND ON” commands in METAL, these SNPs were recoded to ensure consistent allele coding across the samples analyzed. Because the German sample was genotyped on the Illumina platform that contains no palindromic SNPs, we used that sample as the gold standard. A final check identified 36 SNPs whose allele frequency ranges still exceeded 0.2. Most of these were not palindromes, and those that were did not show complementary allele frequencies. We concluded these were unreliable SNPs and dropped them from the analysis.

Selected results were confirmed, and heterogeneity statistics were calculated, using Comprehensive Meta-analysis version 2.0. This performs a random-effects meta-analysis that is robust to sample heterogeneity, as well as explicit tests of heterogeneity using Q and I statistics. In order to account for overlap in control samples, we grouped cases from the NIMH-BP and STEP-BD samples (which do not overlap), and compared them, as a whole, with the controls used in the STEP-BD report, as recommended by some authorities. We used a threshold of genome-wide significance ($7.2 \times 10^{-8}$) derived from a published, genome-wide simulation of common variants in samples of European ancestry. This threshold is more conservative than the value of $1.6 \times 10^{-7}$ that would represent a Bonferroni-corrected $p=0.05$ for the approx. 318,000 markers tested, but is consistent with accepted thresholds for genome-wide significance. Although we set out to find variants shared in common between BP and MDD, we have considered only BP in other studies. The results at rs2251219 would remain significant even if multiplied by 2 to account for this.

Power Analysis was done with Genetic Power Calculator. We assumed a trait prevalence of 2%, minor allele frequency of 20%, an alpha of $7.2 \times 10^{-8}$, and a marker-allele D' value of 0.8.

Gene Expression Analysis

Brain RNA and genomic DNA samples were obtained from the Stanley Medical Research Institute (SMRI). This collection comprises 3 diagnostic groups, each with 35 samples: healthy control, bipolar disorder and schizophrenia. All experiments were done after the specimen code was broken and they were thus unblinded. RNA samples originated from the
dorsolateral prefrontal cortex. Of these, 101 samples provided sufficient RNA for reverse transcription (Transcriptor First Strand cDNA Synthesis kit with oligo-dT priming, Roche Applied Science, Indianapolis, IN) and subsequent real-time PCR analysis. We amplified PBRM1 mRNA with a primer pair common to all known RefSeq transcript variants (Eurofins MWG operon, Huntsville, AL) and a FAM-labeled probe (Roche Applied Science Universal Probe Libraray probe # 41). No known SNPs overlapped with primer or probe sequences (Supplementary Figure 2). For normalization, we used a pre-designed endogenous control assay interrogating PGK1 (Applied Biosystems, Foster City, CA; catalog number 4333765F, FAM-labeled). Reactions were carried out in triplicate in a 384-well LightCycler 480 (Roche Applied Science) in reaction volumes of 8 µL, with 5 ng of reverse-transcribed RNA, 1x final concentration of Roche LightCycler 480 Probes master mix, 450 nM of each primer, and 125 nM of fluorescent probe. assay efficiencies were determined using two-fold serial dilutions of pooled cDNA. Relative expression levels for PBRM1 were calculated using the efficiency-corrected comparative threshold method 28. We used the sample with the median PBRM1 expression level as calibrator and log2-transformed expression levels. One hundred samples were successfully assayed. (As recommended by SMRI, one case sample was omitted due to a known degenerative neurological disorder). Of the remaining 99 samples (31 with bipolar disorder, 34 with schizophrenia, and 33 healthy controls), 97 were from donors of European ancestry. Ninety-six of these samples were successfully genotyped at rs2251219 with standard exonuclease methods (TaqMan, Applied Biosystems, Boston, MA). Data were analyzed by ANOVA (Xlstat 6.0), with diagnosis as the independent, and relative expression level as the dependent, variable. Specific comparisons were performed with the Tukey HSD test. The entire data set has been uploaded to the SMRI data bank.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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Figure 1.
a. Manhattan plot of the meta-analysis results, generated by Haploview 4.0. Physical position is shown along the x-axis, -log (meta-p-value) is shown along the y-axis, and each chromosome is shown in a distinct color. The red guideline indicates the threshold of genome-wide significance ($7.2 \times 10^{-8}$).
b. Detail of the associated region, generated by SNAP 2.0. Physical position and gene annotations (HapMap release 22) are shown along the x-axis, -log (meta-p-value) is shown on the left y-axis, recombination rate (CEU) on the right y-axis.
c. Linkage disequilibrium ($r^2$) as estimated from HapMap 3 phased genotypes, generated by UCSC Genome Browser. Darker red indicates higher values.
Table 1
Descriptive statistics for the samples analyzed. WTCCC: Wellcome-Trust Case Control Consortium bipolar disorder sample; GAIN Depression: Genetic Association Information Network major depressive disorder sample; STEP-BD: STEP-BD bipolar disorder sample; NIMH Bipolar: NIMH bipolar disorder sample; German: German bipolar disorder sample. Counts refer to subjects who passed all quality control filters (see Methods).

| Sample        | Cases | Case Diagnosis                                  | Controls | Platform               |
|---------------|-------|------------------------------------------------|----------|------------------------|
| WTCCC         | 1854  | Bipolar I, bipolar II, schizoaffective bipolar  | 2943     | Affymetrix 500K        |
| GAIN MDD      | 1722  | Major depressive disorder                       | 1774     | Perlegen               |
| STEP-BD       | 1461  | Bipolar I, bipolar II, schizoaffective bipolar  | 2008     | Affymetrix 500K        |
| NIMH Bipolar  | 1001  | Bipolar I, schizoaffective bipolar              | 1033     | Affymetrix 6.0         |
| German        | 645   | Bipolar I                                       | 1310     | Illumina HumanHap 550  |
| **Total**     | 6683  |                                                | 9068     |                        |
### Table 2

Association results for rs2251219 in each sample and in the meta-analysis.

| Sample       | Allele | Frequency (cases) | Frequency (controls) | Allelic Chi-square | Allelic p-value | OR    | N    |
|--------------|--------|-------------------|---------------------|--------------------|----------------|-------|------|
| Discovery Samples |        |                   |                     |                    |                |       |      |
| NIMH BP      | C      | 0.39              | 0.41                | 1.43               | 0.2312         | 0.93  | 2034 |
| German BP    | C      | 0.38              | 0.44                | 9.51               | 0.0020         | 0.81  | 1955 |
| STEP-BD      | C      | 0.38              | 0.40                | 5.67               | 0.0172         | 0.89  | 1461 |
| WTCCC        | C      | 0.36              | 0.40                | 13.68              | 0.0002         | 0.85  | 4797 |
| GAIN MDD     | C      | 0.39              | 0.42                | 4.98               | 0.0256         | 0.90  | 3496 |
| Pooled discovery |      | 0.38              | 0.41                |                    | 1.1 × 10⁻⁸     | 0.87  | 13743 |
| Replication Samples |     |                   |                     |                    |                |       |      |
| GSK-BP       | C      |                   |                     | 0.0023             | 0.57           | 0.84  | 4677 |
| GSK-Lausanne | C      |                   |                     | not available      | 0.3219         | 1.21  | 1349 |
| GSK-Munich   | C      |                   |                     | 0.1227             | 0.79           | 0.79  | 1792 |
| Pooled Replication |   |                   |                     |                    | 0.0012         | 0.84  | 4677 |
| Pooled Discovery and Replication | |                   |                     |                    | 1.7 × 10⁻⁹     | 0.84  | 18420 |