Family-Based SNP Association Study on 8q24 in Bipolar Disorder

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Previous linkage studies have identified chromosome 8q24 as a promising positional candidate region to search for bipolar disorder (BP) susceptibility genes. We, therefore, sought to identify BP susceptibility genes on chromosome 8q24 using a family-based association study of a dense panel of SNPs selected to tag the known common variation across the region of interest. A total of 1,458 SNPs across 16 Mb of 8q24 were examined in 3,512 subjects, 1,954 of whom were affected with BP, from 737 multiplex families. Single-locus tests were carried out with FBAT and Geno-PDT, and multi-locus test were carried out with HBAT and multi-locus Geno-PDT. None of the SNPs were associated with BP in the single-locus tests at a level that exceeded our threshold for study-wide significance ($P < 3.00 \times 10^{-5}$). However, there was consistent evidence at our threshold for the suggestive level ($P < 7.00 \times 10^{-4}$) from both the single locus and multi-locus tests of associations with SNPs in the genes ADCY8, ST3GAL1, and NSE2. Multi-locus analyses suggested joint effects between ADCY8 and ST3GAL1 ($P = 3.00 \times 10^{-4}$), with at least one copy of the “high risk” allele required at both genes for association with BP, consistent with a jointly dominant–dominant model of action. These findings with ADCY8 and ST3GAL1 warrant further investigation in order to confirm the observed associations and their functional significance for BP susceptibility.

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KEY WORDS: bipolar disorder; dominant–dominant model; genetic association; chromosome 8q24

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INTRODUCTION

Linkage studies have implicated chromosome 8q24 as a promising positional candidate regions in bipolar disorder (BP). Two independent genome-wide scans of BP reported findings in the region that met stringent criteria for genome-wide significance established by Lander and Kruglyak [1995]. Cichon et al. [2001] studied 75 BP pedigrees of German, Israeli, and Italian origin and reported a LOD of 3.6 at marker D8S514 (deCODE genetic map: 124.62 cM; Build 35 physical map: 123,811,284–123,811,642). We [McInnis et al., 2003; Avramopoulos et al., 2004] studied 65 pedigrees of European descent and reported a LOD of 3.32 at marker D8S256 (deCODE genetic map: 145.26 cM; Build 35 physical map: 134,513,446–134,513,788). Moreover, McQueen et al. [2005] recently pooled the primary genotype data from 11 BP genome-wide linkage scans (including from our study) and carried out a non-parametric linkage analysis of the combined data. Two regions on 6q21 and 8q24 achieved genome-wide significance in the pooled data analysis.

In a previous study, we genotyped a panel of SNPs in a sub-sample of 583 affected offspring from 258 nuclear families with prior evidence of linkage to the region. The panel included 249 informative SNPs covering a narrowly defined region of interest of 3.4 Mb centered at DSS256, the peak marker in our genome-wide linkage scan. The findings from this association study have been reported elsewhere [Zandi et al., 2007].

In the current study, we expanded our search of the region to cover a larger territory spanning ~16 Mb and including the marker identified in the genome–scan by Cichon et al. [2001]. In addition, we genotyped all 3,512 individuals from 737 extended pedigrees for which we had available DNA in order to extract maximal information for association testing. We report here the initial results of our primary analyses of these data, which we plan to make publicly available.

METHODS

The Sample

This study combined two samples of families. The first sample was collected by our group in the Mood Disorders Research Program at Johns Hopkins University [McInnis et al., 2003]. The families were ascertained by opportunistic...
Family-Based SNP Association Study on 8q24

Genotype Data

Genotyping was done by CIDR using Illumina technology on a BeadLab system with Golden Gate chemistries. A total of 1,536 SNPs from 123.1 to 139.1 Mb (Build 35) on chromosome 8q24 were selected for genotyping. The SNPs were selected using FESTA (Qin et al., 2006), an LD based tagSNP selection program. It uses the same criterion as LDSelect (Carlson et al., 2004), but uses a more efficient algorithm that selects a smaller number of tagSNPs. The tagSNPs were selected using HapMap Phase I data such that all the common SNPs (minor allele frequency [MAF] > 0.05) identified in the HapMap Phase II data that was tagged at an r² > 0.80 by the SNPs genotyped in the current experiment.

Several steps were taken to clean the genotype data. First, we corrected problematic familial relationships that had been identified through previous analyses of microsatellite marker genotype data collected as part of genome-wide linkage scans with these samples. This included the removal of monozygotic twins and reassignment of established incompatible paternities. If the problems could not be corrected, the offending subjects were removed from the dataset. Second, Mendelian inheritance errors were identified using PEDCHECK, and corrected by removing the problematic genotypes according to a pre-specified algorithm established by CIDR. We then rechecked the data for Mendelian errors using Merlin’s PEDSTATS. Finally, we used Merlin’s error routine to examine the data for evidence of unlikely double recombinations.

Statistical Analyses

For our primary analysis, we carried out single-locus tests of association with each individual SNP using two different methods. First, we conducted an allelic test of association using FBAT (Horvath et al., 2001). FBAT is a flexible program that is appropriate for analyses of family data. Under certain conditions, the FBAT reduces to the commonly used transmission/disequilibrium test (TDT). However, FBAT is more general and allows tests of associations that are robust to population confounds in the case where parental data are missing and/or other offspring are included in the analysis (Laird and Lange, 2006). We used the bi-allelic mode and examined the additive model for counting alleles. We specified the option to calculate the variance empirically in order to provide valid tests of association in the presence of linkage (Lake et al., 2000). Second, we conducted genotypic tests of association using the program Geno-PDT (Genotype-Pedigree Disequilibrium Tests) (Martin et al., 2003). Geno-PDT provides valid tests of association in the presence of linkage for genotypes in extended pedigrees. We obtained global 2 degrees of freedom significance values for each test.

We then carried out multi-locus tests of association again using two different methods. First, we tested haplotypes of adjacent SNPs using HBAT (Horvath et al., 2004). It has been shown that in some situations (such as when r² = 1 between the risk variant and a particular multi-SNP haplotype) haplotypes may provide more information for association than corresponding single-locus tests (Clayton et al., 2004). HBAT is an elaboration of FBAT that allows for family-based association tests of haplotypes, even when the phasing of the haplotypes is ambiguous. We used a sliding window approach to test haplotypes of 2, 3, and 4 adjacent SNPs across the region of interest. Significance values were obtained for tests of each specific haplotype. Second, we tested for associations with combinations of genotypes between pairs of SNPs using multi-locus Geno-PDT (van der Walt et al., 2004). This program does not require phase information and, is therefore useful for
dissecting joint effects of SNPs. It provides significance values for individual as well as global tests of the different genotype combinations. To limit the number of tests, we only considered pair-wise combinations of SNPs that provided at least suggestive evidence (defined as below) in the single locus tests.

For all tests of association described above, subjects diagnosed with BPI, schizo-affective disorder, bipolar type (SABP), or bipolar II disorder (BPII) were included as affected. This model of affection was chosen because the best findings on chromosome 8q24 from the previous linkage studies cited above were all obtained under such a model. Subjects who were determined to be never mentally ill were included as unaffected. All other subjects were included with unknown status.

We set the threshold for declaring a finding significant to $P < 3.00 \times 10^{-5}$, reflecting a nominal level of $P < 0.05$ and correcting for 1,458 tests. We used the probability of identifying one false positive out of 1,458 tests ($P < 7.00 \times 10^{-4}$) as the threshold for declaring a finding suggestive. These thresholds do not account for the dependence of the 1,458 SNPs tested or for the multiple methods that were applied to the data.

Fig. 1. Results from the FBAT and Geno-PDT tests for all 1,458 SNPs across the region of interest are shown. Open triangles show the negative log $P$ values for the FBAT test under the additive model with the empirical estimator of the variance to appropriately account for association in the presence of linkage. Open circles show the negative log $P$ values for the Geno-PDT global tests with 2 degrees of freedom. Highlighted are two regions of interest that are shown at greater resolution in Figures 2 and 3.

Fig. 2. Detailed findings from the targeted region around ST3GAL1 including: the results from the single-locus tests with FBAT (open triangles) and Geno-PDT (open circles) of all SNPs in the more narrowly defined region; any findings from the multi-locus haplotype analyses with suggestive evidence ($P < 7.00 \times 10^{-4}$) of association (black bar); a snapshot of the region from UCSC Golden Path showing tracks of known genes and a measure of conservation across 17 species [Siepel et al., 2005]; and a picture of the linkage disequilibrium (LD) structure from Haploview (stronger LD in darker shades of black) with blocks defined by the algorithm of Gabriel et al. [2002]. [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]
RESULTS

The final dataset consisted of 3,512 genotyped subjects, including 1,383 males and 2,129 females. Of these, 1,954 were affected (94 SABP, 1,546 BPI, and 314 BPII), 513 were unaffected (i.e., never mentally ill), and 1,045 had some other diagnosis and were included in the analyses as unknown phenotype. There were a total of 737 families, consisting of 1,142 nuclear families. The nuclear families broke down into 1,840 parent affected offspring triads (573 with zero genotyped parents, 641 with one genotyped parent, and 626 with two genotyped parents) and 333 discordant sib-pairs.

A total of 1,458 SNPs met our quality control standards and were examined across the ~16 Mb region of interest at a mean density of 1 SNP per 11 kb with five gaps of greater than 100 kb.

The results of the single locus tests of association between each SNP and BP using FBAT and Geno-PDT are shown in Figure 1, with greater detail at two sites of particular interest shown in Figures 2 and 3. None of the SNPs that met our quality control standards were associated at a level that exceeded our threshold for study-wide significance ($P < 3.00 \times 10^{-4}$). Data for SNPs that were associated at the suggestive threshold of $P < 7.00 \times 10^{-4}$ on either of the single locus tests are provided in Table I. There were six such SNPs, none of which significantly deviated from HWE. These included rs6986303 (FBAT, $P = 2.82 \times 10^{-4}$) in the fourth intron of ST3GAL1; rs1411187 (FBAT, $6.40 \times 10^{-4}$) and rs3750889 (FBAT, $P = 3.40 \times 10^{-4}$) in the first and second introns, respectively, of ADCYS; and rs7462286 (Geno-PDT, $P = 7.00 \times 10^{-4}$) in the 11th intron of DDEF1. The other two SNPs were not located in genes and included rs10101440 (FBAT, $P = 4.99 \times 10^{-4}$) 29 kb 5' (centromeric) of the nearest gene, NSE2, and rs1668875 (Geno-PDT, $P = 5.00 \times 10^{-4}$) 407 kb 3' (centromeric) of MYC.

To perform multi-locus tests of association, we derived haplotypes by sliding windows of 2, 3, and 4 adjacent SNPs across the region of interest. We observed seven clusters of haplotypes with suggestive evidence of association exceeding the threshold of $P < 7.00 \times 10^{-4}$. Three of these included SNPs that were highlighted in the single locus tests described above. The first set of haplotypes included both rs3750889 and rs1411187 and straddled the second exon of ADCYS ($P = 3.40 \times 10^{-4}$). The second included rs10101440 and spanned the entire gene of NSE2, which only consists of two exons ($P = 3.80 \times 10^{-4}$). The third included rs6986303 and extended from the fourth intron to 2.3 kb past the 3' end of ST3GAL1 ($P = 6.20 \times 10^{-4}$). The other four clusters of haplotypes included SNPs that were not highlighted in the single locus tests. The most significant of these ($P = 3.90 \times 10^{-5}$) was approximately 657 kb away from ST3GAL1 in a relative gene desert that was implicated in an earlier, more limited study of the region (Zandi et al., 2007). The other three were 27 kb 5' (telomeric) of KCNQ3 ($P = 4.30 \times 10^{-4}$), 1.6 Mb 3' (telomeric) of MYC ($P = 5.90 \times 10^{-4}$), and 246 kb 5' (centromeric) of MYC ($P = 3.50 \times 10^{-4}$).

We next carried out an additional multi-locus test of associations with Geno-PDT. There was only one combination of SNPs in which the pairwise test exceeded the threshold of suggestive evidence ($P < 7.00 \times 10^{-4}$). This was between 27 kb 5' of ST3GAL1 and rs1411187 in ADCYS (Table II). The $P$-value for the global pairwise test was 0.0003. Examination of the results suggested that over-transmission to the affected cases (as evidenced by a positive Z-score >1.0) occurred whenever there was at least one copy of the “high-risk” (T) allele at both SNPs.

DISCUSSION

Previous linkage studies indicate chromosome 8q24 is one of the more promising regions to search for BP susceptibility genes (Cichon et al., 2001; McInnis et al., 2003; Avramopoulos et al., 2004; McQueen et al., 2005). We carried out a large, family-based association study with a panel of SNPs spanning ~16 Mb across the locus. We report here the results of our primary analyses of these data. There was consistent evidence

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<th>P-value</th>
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1Results are reported for SNPs that are associated with BP at $P < 7.00 \times 10^{-4}$ with either FBAT or Geno-PDT; the FBAT tests are conducted under the additive model with the empirical variance estimator to appropriately account for association in the presence of linkage, and the Geno-PDT results are shown for the 2 degrees of freedom global test.

2Physical position is based on the May 2004 human reference sequence (Build 34).

3The minor allele is shown.

| Table I. Results From Single SNP Tests With Suggestive Evidence of Association Using FBAT and Geno-PDT | Family-Based SNP Association Study on 8q24 | 615 |
at the suggestive level from both the single locus and multi-locus tests of associations with several SNPs in genes across the region, including ADCY8, ST3GAL1, and NSE2. It is possible there are multiple genes in the region that contribute to BP susceptibility. Indeed, it may have been the clustering of such genes that led to a detectable linkage signal which originally drew attention to the region. The associations with ADCY8 and ST3GAL1 are of particular interest, because of evidence suggesting joint effects between the two genes.

ADCY8 codes for adenyl-cyclase in the brain. This enzyme catalyzes the formation of cyclic AMP (cAMP), an important second messenger that has been implicated in BP and is a target for lithium and other mood stabilizing agents [Mork, 1993; Perez et al., 2000; Stewart et al., 2001]. Thus, it is a promising functional candidate gene for BP. Interestingly, a recent study of another candidate region in BP on chromosome 18p reported an association with a SNP that is 1 Mb away from ADCYAP [Mulle et al., 2007], adenyl cyclase activating polypeptide 1, which stimulates adenyl cyclase in pituitary cells and cAMP levels in target cells.

ST3GAL1 codes for a membrane-bound protein that catalyzes the transfer of sialic acid from CMP-sialic acid to galactose containing substrates. It belongs to a family of glycosyltransferase proteins, several of which have been found to be highly expressed in the developing brain and to play a role in neurogenesis [Angata et al., 2004]. Recently, a member of this protein family, SIAT8B on chromosome 15q26, has been associated with schizophrenia in two separate studies in Japanese [Arai et al., 2006] and Chinese [Tao et al., 2006] populations, providing support for the potential relevance of
TABLE II. Results From Multi-Locus Test Using Geno-PDT With rs1411187 and rs6986303

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Global test*  
\( \chi^2 = 28.90, P = 3.00 \times 10^{-4} \)

Trans, transmitted; NotTrans, not transmitted; Aff, affected sib; Unaff, unaffected sib.

*There is 1 degree of freedom for the genotype specific tests, and 8 degrees of freedom for the global test.

The current study examined a number of important strengths. It examined one of the largest samples of BP families available for genetic studies. Moreover, all members of the families with available DNA were genotyped, yielding maximal information from the data. We conservatively estimated, assuming one affected trio per family and a disease allele frequency of 0.30 under a multiplicative model, that at the level of study wide significance specified above this sample had 80% power to detect a susceptibility locus with a genotypic relative risk as small as 1.50. Finally, the family-based tests of association provided protection against spurious findings due to population stratification. A limitation of the study is that it included families collected under two different protocols, which could have introduced heterogeneity into the combined sample and biased the results toward the null. However, our group was involved in the design and conduct of both collection efforts, and the ascertainment and assessment procedures were comparable. Another limitation of the study is that it was originally designed based on HapMap Phase I data, and as a result, the coverage of the currently known common variation is incomplete.

The findings from the current study suggest several loci on chromosome 8q24 may be associated with BP and warrant closer investigation. We intend to follow-up these findings by genotyping a denser panel of SNPs around the identified loci, especially in the ST3GAL1 and ADCY8 genes, and to fill in the current gaps in coverage. Our goal is to confirm which of the observed associations are real, and ultimately to determine their functional relevance for BP susceptibility. We have made the data from this study publicly available (at http://bioinformooods.jhmi.edu/chr8project/) in order to encourage the development of new methods to analyze complex datasets of this kind and to facilitate continued research on chromosome 8q24 in BP.

ACKNOWLEDGMENTS

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