Assessment of First and Second Degree Relatives of Individuals With Bipolar Disorder Shows Increased Genetic Risk Scores in Both Affected Relatives and Young At-Risk Individuals

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Recent studies have revealed the polygenic nature of bipolar disorder (BP), and identified common risk variants associated with illness. However, the role of common polygenic risk in multiplex families has not previously been examined. The present study examined 249 European-ancestry families from the NIMH Genetics Initiative sample, comparing subjects with narrowly defined BP (excluding bipolar II and recurrent unipolar depression; n = 601) and their adult relatives without BP (n = 695). Unrelated adult controls (n = 266) were from the NIMH TGEN control dataset. We also examined a prospective cohort of young (12–30 years) offspring and siblings of individuals with BPI and BPII disorder (at risk; n = 367) and psychiatrically screened controls (n = 229), ascertained from five sites in the US and Australia and assessed with standardized clinical protocols. Thirty-two disease-associated SNPs from the PGC-BP Working Group report (2011) were genotyped and additive polygenic risk scores (PRS) derived. We show increased PRS in adult cases compared to unrelated controls (P = 3.4 × 10⁻⁵, AUC = 0.60). In families with a high-polygenic load (PRS score ≥32 in two or more subjects), PRS distinguished cases with BPI/BPII from other relatives (P = 0.014, RR = 1.32). Secondly, a higher PRS was observed in at-risk youth, regardless of affected status, compared to unrelated controls (GEE-χ² = 5.15, P = 0.012). This report is the first to explore common polygenic risk in multiplex families, albeit using only a small number of

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robustly associated risk variants. We show that individuals with BP have a higher load of common disease-associated variants than unrelated controls and first-degree relatives, and illustrate the potential utility of PRS assessment in a family context.

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INTRODUCTION

Large scale collaborative genome-wide association studies (GWAS) have identified a number of risk loci significantly associated with bipolar disorder (BP), including ODZ4 [Sklar et al., 2011; Muhleisen et al., 2014], CACNA1C [Ferreira et al., 2008; Sklar et al., 2008], ANK3 [Ferreira et al., 2008; Schulze et al., 2009; Muhleisen et al., 2014], NCAN [Cichon et al., 2011], C15orf53 [Ferreira et al., 2008], and DGKH [Baum et al., 2008a,b]. Individually, each of those genes/loci contributes only a small fraction toward overall disease risk, typically <1% of the phenotypic variance [Ferreira et al., 2008]. It is now understood that multiple genes containing both common and rare variants contribute to the genetic architecture of BP [Sullivan et al., 2012], and there are significant overlaps in the single nucleotide polymorphism (SNP)-based heritabilities of BP with both schizophrenia and major depression [Lee et al., 2013]. Indeed, variation across many thousands of common risk variants together (termed polygenic risk [Purcell et al., 2009]) contributes a substantial proportion (i.e., 25–40%) of the percentage of phenotypic variance at a population level [Lee et al., 2011; 2013]—although most of those loci do not individually reach genome-wide significance thresholds for disease association with current sample sizes [Craddock and Sklar, 2013; Dudbridge, 2013].

Examination of the cumulative effects of inheriting multiple risk alleles—each of which are significantly or nominally associated with disease risk—has been able to powerfully differentiate groups of cases from controls in independent population-based studies [Purcell et al., 2009, 2014; Patel et al., 2010; Ayalew et al., 2012; Terwisscha van Scheltinga et al., 2012]. However, despite phenotypic aggregation within families [McGuffin et al., 2003; Lichtenstein et al., 2009], no studies have so far examined polygenic risk incorporating these common genetic factors in a family context in adults, with only one group to date reporting on polygenic risk in adolescent offspring of individuals with BP [Whalley et al., 2012, 2013].

While first-degree relatives of individuals affected with BP would be expected on a theoretical basis to have a higher load of specific disease-associated risk alleles than control individuals, this has not previously been examined empirically. This is primarily because, until recently, we have had very little knowledge about the specific DNA variants that contribute to risk for BP. However, as sample sizes are steadily increasing through the Psychiatric Genomics Consortium (PGC), and our power to detect such risk variants on a population level is improving, we are gaining a greater understanding of the underlying genetic contributors leading to BP. In 2011, Sklar et al. identified 38 variants (pared to 34 by assessment of independence with linkage disequilibrium) that contribute to disease risk $(P < 5 \times 10^{-5})$ using a discovery sample of 7,481 individuals with BP and 9,250 controls. These SNPs were replicated in an independent cohort comprising 4,496 cases and 42,422 controls, with more SNPs than expected showing $P < 0.01$, $P < 0.05$, and the same direction of effect [Sklar et al., 2011]. Hence, a substantial number of these SNPs may be expected to represent true risk variants (or markers for such alleles) for bipolar disorder. Here, we examined polygenic risk derived from those specific variants in first degree relatives of individuals with BP.

The present study tested the hypothesis that first-degree relatives who are affected with BP would have a higher polygenic risk load of common risk variants in comparison to their relatives without BP, and that individuals at increased familial risk of BP would have a higher polygenic load of risk variants than control individuals. We tested this hypothesis using three cohorts: (1) a group of singleton cases with BP and controls from the National Institute of Mental Health (NIMH); (2) a family cohort of adult relatives of individuals with and without diagnoses of BP from the NIMH Bipolar Genetics Initiative; and (3) a prospective cohort of adolescents and young adults who are at increased familial risk of developing BP due to the presence of a first degree relative with a diagnosis of BP (henceforth termed “at-risk”) and controls.

MATERIALS AND METHODS

Adult Participants From Family Studies

Two datasets were employed. The first comprised adult subjects of European ancestry $(n = 1,947)$ drawn from the NIMH Genetics Initiative bipolar disorder family samples (waves I–IV European American families, $n = 249$ families, average of 6–7 subjects across three generations per family) [Nurnberger et al., 1997; Smith et al., 2009]. Subject diagnoses were obtained via standard best estimate (BEFD) procedure (details in supplementary information), and were diagnosed as having bipolar disorder type I (BPI, $n = 561$); schizoaffective disorder-bipolar type (SAB, $n = 40$); bipolar disorder type II (BPII, $n = 119$); recurrent unipolar depressive disorder (UPR, $n = 155$); or single episode unipolar depressive disorder (UPS, $n = 107$). Primary analysis utilized a narrow definition of case status, whereby subjects were

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defined as affected if having received a diagnosis of BPI or SAB (n = 601). Subjects with BPII and UPR (n = 274) were excluded on the basis of unknown overlapping etiology, as were relatives with unknown diagnosis (n = 85). All other relatives (n = 695) included those with other mental disorders ([i.e., those with any DSM diagnosis excluding BPI, SAB, BPII, or UPR, n = 292], never mentally ill [n = 296], or with single episode depression [n = 107]), and were analyzed as a single group. Due to the small numbers of relatives with no psychiatric diagnoses (i.e., only one per family) comparisons between cases and never mentally ill relatives were not attempted.

The comparison dataset comprised unrelated adult controls of European origin drawn from the NIMH TGEN Control dataset subjects (n = 403) from the control collection of Sanders et al. [2010], who were screened to exclude those with any major mood disorder or psychosis (https://www.tgen.org/). DNA was extracted from whole blood by the Rutgers University Cell and DNA Repository (US participants) or Genetic Repositories (Australian participants). The 596 participants came from five independent sites in the US [Nurnberger et al., 2011] (Johns Hopkins University; University of Michigan; Washington University in St. Louis; Indiana University) and Australia [Roberts et al., 2013] (University of New South Wales). "At-risk" subjects were recruited from families who had previously participated in BP genetics studies [Nurnberger et al., 1997; Dick et al., 2003; Fullerton et al., 2010], a specialized BP research clinic [Mitchell et al., 2009], were referred by clinicians or mental health consumer organizations, or responded to other forms of publicity. Subjects were recruited using a "top-down" ascertainment method: that is, all offspring of a proband with a confirmed DSM-IV diagnosis of bipolar disorder type I (BPI), type II (BPII), or schizoaffective disorder bipolar-type (SAB) who were in the age range 12–30 were eligible for inclusion, independent of the diagnostic status of the offspring. The participants were predominantly (94%) children or siblings of a proband; the remaining 6% comprised second-degree relatives ascertained from families with multiple cases of BP. Control participants were in the same age range, but had no first-degree relative (parent or sibling) with a DSM-IV diagnosis of BPI or BPII, recurrent major depression (UPR), schizophrenia, recurrent substance abuse, or any past psychiatric hospitalization; and no parent with a first-degree relative who had a past mood disorder hospitalization or history of psychosis. Control subjects were recruited via general medical clinics, motor vehicle records, print and electronic media, and notice boards in universities and local communities.

Peripheral blood samples were collected from a total of 367 at-risk and 229 control individuals for genetic analysis. DNA was extracted from whole blood by the Rutgers University Cell and DNA Repository (US participants) or Genetic Repositories Australia (Australian participants). The 596 participants came from 426 families: the majority being single offspring (n = 299); although 127 families with ≥2 offspring were included (90 with 2 offspring, 30 with 3 offspring, 5 with 4 offspring, and 2 with 5 offspring).

Institutional Review Board Approval
Written informed consent (or assent with parental consent for subjects <18 years old) was obtained from all participants after a thorough explanation of the study. All protocols were approved through the individual hospital and University ethics committees (Institutional Review Boards) at each of the participating US university medical centers, University of New South Wales Human Research Ethics Committee, and the South Eastern Sydney Illawarra Area Health Service, Australia.

Ascertainment of Clinical Diagnoses and Demographic Information
Methods for ascertainment and diagnoses for the NIMH Genetics Initiative family dataset have been described extensively elsewhere [Nurnberger et al., 1997; Smith et al., 2009] and are summarized in supplementary information.

For the at-risk cohort, proband consensus DSM-IV diagnoses were determined by two independent psychiatrists using best estimate methodology [Leckman et al., 1982], using information from an adapted version of the Schedule for Affective Disorders and Schizophrenia for School-Age Children—Present and Lifetime Version (K-SADS-BP) [Kauffman et al., 1997; Geller et al., 2001], the Diagnostic Interview for Genetic Studies (DIGS) Version 4 [Nurnberger et al., 1994], the Family Interview for Genetic Studies (FIGS) [Maxwell, 1992], and medical records (where available). For participants under the age of 22, the K-SADS-BP was administered as the diagnostic instrument. The FIGS was administered to all participants 18 years of age or older, with parents completing FIGS for participants under 18. Ethnicity was assessed by self-report via grandparental origin information. The at-risk cohort was mainly of European ancestry (73%), with subjects of Asian and African ancestry each accounting for less than 10% of the sample (Table I).

Marker Selection, Genotyping, and Quality Control
We chose for genotyping 38 SNPs that were robustly implicated in bipolar disorder risk, on the basis of prior evidence of genetic association (P < 5 × 10⁻⁵) from the PGC [Sklar et al., 2011]; two (rs3968, rs8006348) failed assay design.

The remaining 36 SNPs [Sklar et al., 2011] were genotyped at two sites, using iPLEX GOLD chemistry on the Sequenom MassArray (Supporting Information Table S1). Tests for Hardy–Weinberg equilibrium, linkage disequilibrium, genotype missingness and allelic, and genotypic frequency comparisons were conducted using PLINK [Purcell et al., 2007]. Three failed genotyping or Hardy–Weinberg equilibrium (P < 0.001) in either the US or Australian at-risk sample (rs12912251, rs4332037, rs7578035), and one was in linkage disequilibrium with another SNP (rs11168751 with rs2070615; r² = 0.141, D' = 1.0). These SNPs were excluded from polygenic risk score analysis, leaving 32 SNPs for determination of polygenic risk score. The successfully genotyped SNPs had a >99.6% genotype pass rate.

NIMH controls were genotyped on the Affymetrix 6.0 array. Imputation was employed via IMPUTE v0.5.0 [Marchini et al.,...
Polygenic Risk Score Determination

To determine the additive polygenic load in each subject, a score of one was given for each risk allele carried, so an individual subject’s score could range from 0 to 64. Weighted-additive polygenic risk scores were computed using the score function in PLINK [Purcell et al., 2007]. Scores were weighted either by: (1) the effect size of each SNP, as determined by the odds ratio of the risk allele from the original discovery GWAS study by Sklar et al. [2011] for Europeans (±0.022–0.032), and were similar for both directly observed and imputed SNPs (±0.031 and ±0.025, respectively) (Supporting Information Table S1).

Statistical Analysis

To test whether cases or at-risk individuals within families had a higher average polygenic risk score as compared with controls, a linear Generalized Estimating Equations (GEE) model was used, with a one-sided test for significance. The GEE model corrects for non-independence of measurement between family members. A one-sided t-test was utilized to test group differences in unrelated subjects. Relative risk calculations were conducted relative to the mean polygenic risk value for the case (bipolar disorder or at-risk) group. Statistical analysis was conducted in SPSS (Version 20.0, IBM Corporation, Armonk, NY). Estimates of the proportion of genetic variance accounted for by the score were calculated via genRoc (http://glimmer.rstudio.com/kn3in/genRoc/) [Wray et al., 2010] and GPRS software (https://gprs.shinyapps.io/start/) [Dudbridge, 2013].

RESULTS

Using polygenic risk scores derived from 32 of the most robustly associated SNPs from the PGC, we conducted a risk load analysis to determine whether a polygenic risk score derived from disease-associated SNPs would: (1) distinguish unrelated BP cases from unrelated control individuals; (2) distinguish relatives affected with BP from their non-BP-affected or “unaffected” adult relatives; and (3) distinguish a young at-risk population from a group of unrelated young controls.

Polygenic Risk Load Analysis in Adult Cases

We used European ancestry subjects from 249 bipolar pedigrees from the NIMH genetics initiative sample. We selected a single case with BPI or SAB from each of the families (n = 236; typically the first subject recruited with a diagnosis of BPI or SAB for each family for whom the maximum number of SNPs were successfully genotyped) and compared polygenic risk scores to unrelated controls from the NIMH control dataset (n = 266).

We found that the average polygenic risk score was higher for singleton BPI or SAB cases than unrelated controls [mean = 32.67 ± 3.85 and 31.35 ± 3.90, respectively; t(1,501) = −3.81, P = 7.08 E-05, OR = 1.88 (95%CI = 1.31–2.69)] and that a score of ≥32 risk alleles (defined by mean of the case group) was associated with an increased relative risk of BP diagnosis of 1.31 (95%CI = 1.12–1.53).
The proportion of the known genetic variance explained by a genomic profile explains a quarter of the known genetic variance [Wray et al., 2010]. The area under the receiver-operator characteristic curve (AUC) was modestly increased from the null at $P = 0.0088$ and odds ratio $(OR = 3.74)$, with an absolute mean score difference $(ABS[sib_1 - sib_2])$ of $±3.74$ risk alleles (Fig. 2). Less predictably, phenotypically concordant case pairs $(n = 108)$ did not have significantly different absolute mean score differences compared to phenotypically discordant sibpairs $(n = 103)$ and discordant non-case pairs $(n = 95)$ (ANOVA, $F = 1.38, P = 0.25$).

Next, we examined polygenic risk score differences amongst all relatives in a within-family analysis $(n = 249$ families), comparing all cases affected with BPI or SAB $(n = 600)$ to all other relatives $(n = 695)$. We found no significant differences by diagnostic group across all families $(mean = 32.69 ± 3.94)$ vs. $32.45 ± 3.85$; Wald $t = 0.93, P = 0.35$.

**TABLE II. Summary of Area Under the Receiver-Operator Characteristic Curve (AUC), With Incremental Increase of SNP Content in Polygenic Risk Score**

| Number of SNPs | AUC (95%CI) | $P$-value |
|----------------|-------------|-----------|
| 10             | 0.528 (0.477–0.578) | 0.287     |
| 13             | 0.548 (0.498–0.599) | 0.062     |
| 14             | 0.554 (0.504–0.605) | 0.037     |
| 15             | 0.556 (0.506–0.607) | 0.029     |
| 20             | 0.573 (0.523–0.624) | 0.0045    |
| 25             | 0.597 (0.547–0.646) | 1.85E-04  |
| 30             | 0.606 (0.557–0.655) | 4.24E-05  |
| 32             | 0.601 (0.552–0.651) | 9.13E-05  |

SNPs were included in the polygenic risk score in order of decreasing odds ratio from the primary GWAS reported by Sklar et al. [2011], and polygenic risk scores weighted by the odds ratio for each SNP. Mean AUC estimates are given, along with the 95% confidence interval of each measure.

It must be noted however, that while significant, the predictive capacity of the PRS is low, indicative of the small effect sizes of individual risk alleles.

**Polygenic Risk Load Analysis Within Families, Comparing Adult Relatives With Bipolar Disorder to Other Relatives**

We next sought to examine polygenic risk scores within multiplex families, to determine if the average polygenic risk score would be higher in cases compared to other relatives. We used European ancestry subjects from 249 bipolar pedigrees from the NIMH genetics initiative, and applied a narrow diagnostic model, where only subjects with diagnoses of BPI or SAB were considered to be cases $(n = 601)$. The category “all other relatives” $(n = 695)$ included subjects with other mental illness (i.e., those with any DSM diagnosis excluding BPI, SAB, BPII, or UPR) and those who were never mentally ill.

As inheritance of polygenic risk has not previously been examined in a family context, we first examined the relationship between risk scores amongst related individuals in each family. We selected all available sibling pairs from the 249 NIMH families $(n = 777$ pairs), and examined the sibship correlation in polygenic risk score. As expected, risk scores amongst all sibings regardless of phenotype were significantly correlated (Pearson $R = 0.53$, $P = 3.60 × 10^{-57}$), with an absolute mean score difference $(ABS[sib_1 - sib_2])$ of $±3.74$ risk alleles (Fig. 2). Less predictably, phenotypically concordant case pairs $(n = 108)$ did not have significantly different absolute mean score differences compared to phenotypically discordant sibpairs $(n = 103)$ and discordant non-case pairs $(n = 95)$ (ANOVA, $F = 1.38, P = 0.25$).

Next, we examined polygenic risk score differences amongst all relatives in a within-family analysis $(n = 249$ families), comparing all cases affected with BPI or SAB $(n = 600)$ to all other relatives $(n = 695)$. We found no significant differences by diagnostic group across all families $(mean = 32.69 ± 3.94)$ vs. $32.45 ± 3.85$; Wald

![Fig. 1. Risk allele score distribution comparing unrelated controls and singleton cases from NIMH families. Single cases with BPI or SAB $(n = 236$, grey bars) were selected from NIMH families and compared to unrelated controls from the TGEN dataset $(n = 266$, white bars). The relative risk $RR = 1.31$ (95%CI: 1.12–1.53) was calculated with respect to a risk score of $≥32$, with an overall odds ratio of $1.88$ (95%CI: 1.28–2.70). For the purposes of graphical representation, the frequency of risk scores are represented in even integers, and represent bins which include both odd and even scores (i.e., bin 22 is the sum of the frequency of 22 and 23 risk alleles).](image-url)
\[ \chi^2 = 1.19, \quad P = 0.13; \quad RR = 1.06 \ (95\% CI = 0.96–1.17), \quad P = 0.14 \] (Fig. 3A).

However, given the heterogeneity of bipolar disorder we hypothesized that some families will have a higher load of these specific common variants than others, and that polygenic risk score differences may be informative of diagnosis only in families which carry a high load of these specific common variants of small effect, as opposed to families whose illness may be caused largely by inheritance of rare pathogenic variation in a smaller number of key genes.

FIG. 2. Relationship between polygenic risk scores in sibling pairs from NIMH families. A) Distribution of polygenic risk score difference between sibpairs \(n = 777\). B) A significant correlation between polygenic risk scores of sibling pairs was observed \((n = 777,\) Pearson \(R = 0.53, \quad P = 3.60E-57)\).

FIG. 3. Analysis polygenic score by diagnosis in NIMH families comparing cases to all other relatives. The distribution of risk scores in patients affected with BPI or SAB [dark grey bars] were compared to all other relatives [light grey bars]. For the purposes of graphical representation, the frequency of risk scores are represented in even integers, and represent bins which include both odd and even scores [i.e., bin 22 is the sum of the frequency of 22 and 23 risk alleles]. Relative risk (RR) estimates were calculated with respect to an additive risk score of \(\geq 32\). A) All NIMH families \(n = 249\) included 600 cases and 715 other relatives. No significant differences were observed between mean risk scores in case versus all other relative groups \([\text{mean} = 32.69 \pm 3.94 \text{ vs. } 32.45 \pm 3.85; \text{GEE-Wald } \chi^2 = 1.19, df = 1, \quad P = 0.13]\). Relative risk was not significant \([\text{RR} = 1.06 \ (95\% CI = 0.96–1.17), \quad P = 0.14]\). B) Selected NIMH families with a high polygenic load \(n = 202\), where two or more individuals from each family had a risk score of \(\geq 32\). The mean risk score was significantly higher in cases \(n = 518\) compared to all other relatives \(n = 613\) \([\text{mean} = 33.40 \pm 3.62 \text{ vs. } 32.95 \pm 3.70; \text{GEE-Wald } \chi^2 = 4.78, P = 0.014]\). The distribution of risk scores in patients affected with BPI or SAB were shifted significantly towards the right compared to their other relatives, with a significant increase in relative risk in cases \([\text{RR} = 1.32 \ (95\% CI = 1.03–1.70), \quad P = 0.018]\).
Hence, we selected only families in which two or more individuals (regardless of diagnosis) had a high-polygenic risk score (i.e., \( P < 0.05 \)) and repeated the analysis in those 202 families. We found that the polygenic risk score significantly differentiated the diagnostic groups in families with a high-common variant load, such that cases \(( n = 518 \) vs. \( n = 613 \)) had higher polygenic risk scores than other relatives (excluding those with diagnoses of BPII or UPR; \( n = 613 \)) (Wald \( \chi^2 = 4.78, P = 0.014; RR = 1.32 [95\% CI = 1.03–1.70], P = 0.018 \)) (Fig. 3B). This was also significant when subjects affected with BPII were included as cases in the model \(( n = 621 \) cases vs. \( n = 613 \) other relatives; mean \( \chi^2 = 33.31 \pm 3.61 \) vs. \( 32.92 \pm 3.70 \); Wald \( \chi^2 = 3.53, P = 0.030; RR = 1.26 [95\% CI = 1.00–1.60], P = 0.032 \)).

**Polygenic Risk Load Analysis in Young First Degree Relatives At-Risk of Bipolar Disorder**

Next, we examined polygenic risk in young at-risk subjects of European ancestry \(( n = 334 \) vs. 142). The at-risk group showed higher mean scores than controls (GEE-Wald \( \chi^2 = 5.15, P = 0.012 \)) (Fig. 4A). The relative risk estimate for subjects with scores \( \geq 32 \) was 1.20 ([95%CI = 0.99–1.45], \( P = 0.036 \)) (Fig. 4B). Results were similar in analyses weighted by allele frequency (GEE-Wald \( \chi^2 = 5.55, P = 0.009 \)) and odds ratio (GEE-Wald \( \chi^2 = 4.18, P = 0.020 \)).

Expanding the cohort to include subjects from the three main ethnic groups (i.e., European, Asian, and African) in an ethnicity-specific weighted analysis showed an enrichment of risk alleles in at-risk subjects compared to controls (GEE-Wald \( \chi^2 = 3.62, P = 0.029; RR = 1.14 [95\% CI = 1.01–1.29], P = 0.017 \) (Supplementary Information Figure S1)).

**DISCUSSION**

Many genomic variants together contribute to overall risk (termed polygenic risk) for a number of complex traits [Peterson et al., 2011; Hamshere et al., 2013; Meyers et al., 2013], and this genetic architecture is evident in a number of psychiatric conditions—including BP [Purcell et al., 2009; Lee et al., 2012, 2013; Smoller et al., 2013; Bramon et al., 2014]. While the elucidation of the genetic causes for BP has been challenging, the field is progressing in understanding the genetic architecture of this complex disorder (reviewed in [Craddock and Sklar, 2013]) and in identifying specific genes which increase risk [Sklar et al., 2011]. Polygenic risk scores based on multiple genetic variants across the genome [Purcell et al., 2009] have successfully discriminated between groups of unrelated cases and controls [Patel et al., 2010] and also individuals with BP broadly defined as schizoaffective or non-schizoaffective [Hamshere et al., 2011], indicating the potential utility of risk score analysis in clinical diagnoses at a population level. However, it is unclear as to whether polygenic risk score analysis, with or without other clinical or biomarker data, could be useful for diagnosis or risk prediction in persons with a significant family history of BP, particularly given the non-random inheritance of population risk alleles in related individuals and confounding shared environmental effects within a family. This question is also pertinent due to the potential for high rates of sporadic illness in typical gene discovery studies [Yang et al., 2010] which are used to define common polygenic risk. The present
study is the first to examine these common risk factors in the context of inheritance within the family of a BP proband, exploring both adult relatives of known diagnosis, and adolescent or young adult relatives who are at-risk of future BP on the basis of a positive family history.

Determination of polygenic risk scores can be performed using strict (e.g., genome-wide significant P < 5 × 10⁻⁵) or increasingly permissive P-value thresholds to include variation with very small genetic effects which may impact disease status. We chose to focus our study on 32 SNPs that were the most significant independently associated variants in the PGC-GWAS analysis [Sklar et al., 2011]. While we acknowledge that our limited SNP selection represents only a small fraction of the total variation that contributes to bipolar disorder risk [Lee et al., 2011, 2013], the selected SNPs arguably represent the largest effect sizes on a population level and are potentially least subject to statistical fluctuation and type I error.

We first sought to determine whether this SNP panel could be useful in distinguishing individuals with a diagnosis of narrowly defined bipolar disorder from unrelated control individuals. The AUC value of the SNP panel (0.60) was small but significant, and slightly higher than the median AUC values based on “known” bipolar risk variants (n = 3) with genome-wide SNP data at a more liberal P-value inclusion threshold of P < 0.0001–0.01 that was previously reported in the WTCCC dataset [Evans et al., 2009]. This may indicate that the larger PGC analysis has a greater signal-to-noise ratio in the top risk SNPs than those identified in the earlier WTCCC analysis, and is consistent with simulation studies which indicate that the accuracy of diagnostic prediction using polygenic risk scores depends on the size of the training sample [Purcell et al., 2009; Dudbridge, 2013].

We found that polygenic risk scores significantly distinguished between diagnostic groups in families with a higher polygenic risk load, but not in families with moderate–low polygenic loads—consistent with genetic heterogeneity across families. Risk scores were not sufficiently specific to classify diagnostic status on an individual basis. Remarkably, we were also able to identify group differences between young individuals at-risk of BP and young controls.

The PGC discovery sample from which the selected SNPs were derived was ethnically European, and as such we restricted our primary analysis to subjects of European descent. By expanding the analysis to include the three major ethnic groups represented in the at-risk cohort, we assume the same SNPs and alleles will confer risk across different ethnic groups. There is little clear evidence regarding ethnic-specific locus-heterogeneity, partly due to the smaller sample sizes currently available for non-European gene discovery studies. However, a recent meta-analysis of European and Asian GWAS data has shown that Asian subjects tend to have the same direction of effect for the most significantly associated loci (P < 1e-06) with only ~2% of the top 41 SNPs showing a different direction of effect to the European samples [Chen et al., 2013]. Additional method development to address the use of polygene scores in the presence of ethnic differences would be useful.

The purpose of our study was to determine whether a small number of common risk variants could be useful in distinguishing relatives of BP probands who are also ill, or may become ill in the future. Despite the close genetic relationship between extended family members, and a non-random inheritance of risk alleles within a family, we were able to show that the polygenic risk score did serve as a marker of ill versus well relatives on a group basis. We would not recommend clinical application of such a score at this time, but additional implementation of similar methods in longitudinal clinical research studies is certainly called for.

Only one other group thus far has examined polygenic risk in a prospectively recruited bipolar cohort, showing an increased genome-wide polygenic risk score in 70 at-risk individuals compared to 60 controls [Whalley et al., 2013] at baseline assessment. This group has also reported polygene associations with limbic brain activation during functional MRI [Whalley et al., 2012], and white matter integrity measures from DTI in a bipolar at-risk cohort [Whalley et al., 2012]. Neuroimaging biomarkers have previously been identified in the Australian at-risk subjects which were part of the current study, with a lack of recruitment of the inferior frontal gyrus in the high-risk participants compared to healthy controls during an fMRI emotion inhibition task [Roberts et al., 2013]. Further investigations on the impact of polygenic risk on neuroimaging biomarkers may be particularly informative.

While this paper describes baseline associations of this at-risk cohort, future investigation of this sample—which is being prospectively evaluated as subjects transition through the peak period of risk for the development of BP—will be able to determine if polygenic risk scores are useful in future risk prediction for the development of bipolar disorder. It remains to be seen as to whether polygenic risk scores alone [Patel et al., 2010], or in conjunction with early clinical signs, exposure to psychosocial risk factors, and other potential biomarkers [Brietzke et al., 2012] may provide a more robust predictor of future illness. The predictive power of polygenic risk scores is likely to improve with increased discovery sample sizes and the assessment of a larger number of both common and rare genetic variants within the prediction models [Sullivan et al., 2012; Chatterjee et al., 2013; Craddock and Sklar, 2013; Dudbridge, 2013].

LIMITATIONS

Some of the parents or relatives of the US at-risk subjects were ascertained from the NIMH families which were included in the PGC discovery sample, and hence our family samples are not entirely independent. However, there is no overlap between the Australian at-risk subjects and the PGC discovery sample, nor are the Australian at-risk subjects related to this sample. While the overall contribution of those NIMH relatives to the PGC discovery sample was very small (<2% of the cases from the PGC discovery dataset), our findings should be considered as an extension rather than an independent replication of the PGC findings (see also discussion in [Wray et al., 2013]). Using only the most significant SNPs is also a limitation of our study, and we acknowledge that many more variants of importance in conferring risk to bipolar disorder have not been assessed. Conversely, this is also a strength of our study, as we were able to directly genotype each variant in the at-risk and family samples, rather than relying on imputed data or surrogate SNPs. Indeed, fewer than half of the 32 SNPs in our panel are represented on any one high-density SNP chip currently commercially available, although direct genotyping of a larger number of SNPs showing nominally significant association will
be possible with the PsychChip (Illumina, San Diego, CA). Imputation was used in the determination of genotypes in the unrelated control group for AUC estimation, and while imputation accuracy was high (97.4% concordance), this is a limitation of the AUC analysis and, together with the small overlap between the PGC discovery sample and the US family samples, should be taken into consideration with interpretation of the AUC data [Wray et al., 2013]. It should be noted that limitations of genotyping platform and imputation did not apply to the key within-family results. We also applied a genetic model which assumes additivity of risk alleles in a single aggregate score as per the methods used by the PGC [Purcell et al., 2009], although it should be noted that this is a simplistic model which does not account for possible multiplicative interaction effects between genes or genes and environment.

CONCLUSIONS

Our study provides conceptual support to the notion that polygenic risk scores may be useful in prospective risk prediction for bipolar disorder. This may lead to future opportunities for early identification and intervention strategies, such as prophylactic pharmaceutical treatment, environmental modifications, or targeted psychological interventions (reviewed in [McMurrich et al., 2012; McNamara et al., 2012]) to reduce the impact and development of symptoms, improve quality of life, and long-term outcomes for patients.

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We have used DNA samples and clinical data from the National Institute of Mental Health Genetics Initiative bipolar disorder family sample (https://nimhgenetics.org/available_data/bipolar_disorder) which is housed at the NHGRI Repository (http://www.genome.gov/19518664), and thank all those involved in generating this resource, who are listed in the Supplementary Material. Genome-wide SNP genotyping of the NIMH samples was performed through the Genetic Association Information Network under the direction of the Bipolar Genome Study (BiGS) Consortium. We thank Kerrie Pierce (NeuRA) and Tamara McDonald (AGRF) for their assistance with sample preparation and genotyping.

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Control subjects from the National Institute of Mental Health Schizophrenia Genetics Initiative (NIMH-GI), data and biomaterials are being collected by the “Molecular Genetics of Schizophrenia II” (MGS-2) collaboration. The investigators and co-investigators are: ENH/Northwestern University, Evanston, IL, MH059571, Pablo V. Gejman, M.D. (Collaboration Coordinator; PI), Alan R. Sanders, M.D.; Emory University School of Medicine, Atlanta, GA, MH59587, Farooq Amin, M.D. (PI); Louisiana State University Health Sciences Center; New Orleans, Louisiana, MH067257, Nancy Buccola A.P.R.N., B.C., M.S.N. (PI); University of California-Irvine, Irvine, CA, MH60870, William Byerley, M.D. (PI); Washington University, St. Louis, MO, U01, MH60879, C. Robert Cloninger, M.D. (PI); University of Iowa, Iowa, IA, MH59566, Raymond Crowe, M.D. (PI), Donald Black, M.D.; University of Colorado, Denver, CO, MH059565, Robert Freedman, M.D. (PI); University of Pennsylvania, Philadelphia, PA, MH598657, Douglas Levinson, M.D. (PI); University of Queensland, Queensland, Australia, MH059588, Bryan Mowry, M.D. (PI); Mt. Sinai School of Medicine, New York, NY, MH59586, Jeremy Silverman, Ph.D. (PI).

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SUPPORTING INFORMATION

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