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#### 1 Genomic dissection of bipolar disorder and schizophrenia including 28 subphenotypes

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#### 100 Summary

101 Schizophrenia and bipolar disorder are two distinct diagnoses that share symptomology. 102 Understanding the genetic factors contributing to the shared and disorder-specific symptoms will 103 be crucial for improving diagnosis and treatment. In genetic data consisting of 53,555 cases 104 (20,129 BD, 33,426 SCZ) and 54,065 controls, we identified 114 genome-wide significant loci 105 implicating synaptic and neuronal pathways shared between disorders. Comparing SCZ to BD 106 (23,585 SCZ, 15,270 BD) identified four genomic regions including one with disorder-107 independent causal variants and potassium ion response genes as contributing to differences in 108 biology between the disorders. Polygenic risk score (PRS) analyses identified several significant 109 correlations within case-only phenotypes including SCZ PRS with psychotic features and age of 110 onset in BD. For the first time, we discover specific loci that distinguish between BD and SCZ and 111 identify polygenic components underlying multiple symptom dimensions. These results point to 112 the utility of genetics to inform symptomology and potentially treatment.

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#### 115 Introduction

Bipolar disorder (BD) and schizophrenia (SCZ) are severe psychiatric disorders and among the leading causes of disability worldwide(Whiteford et al., 2013). Both disorders have significant genetic components with heritability estimates ranging from 60-80%(Nöthen et al., 2010). Recent genetic and epidemiological studies have demonstrated substantial overlap between these two disorders with a genetic correlation from common variation near 0.6-0.7(Cross-Disorder Group of the Psychiatric Genomics Consortium, 2013) and high relative risks (RR) among relatives of both BD and SCZ patients (RRs for parent/offspring: BD/BD: 6.4, BD/SCZ: 2.4; SCZ/BD: 5.2, 123 SCZ/SCZ: 9.9)(Lichtenstein et al., 2009). Despite shared genetics and symptomology, the current 124 diagnostic systems("Diagnostic and Statistical Manual of Mental Disorders | DSM Library," 125 n.d.) ("WHO | International Classification of Diseases," n.d.) adhere to historical distinctions from the late 19<sup>th</sup> century and represent BD and SCZ as independent categorical entities differentiated 126 127 on the basis of their clinical presentation, with BD characterized by predominant mood symptoms, 128 mood-congruent delusions and an episodic disease course and SCZ considered a prototypical 129 psychotic disorder. Identifying genetic components contributing to both disorders provides insight 130 into the biology underlying the shared symptoms of the disorders.

131 While the shared genetic component is substantial, studies to date have also implicated genetic 132 architecture differences between these two disorders(Curtis et al., 2011; Ruderfer et al., 2014). A 133 polygenic risk score created from a case only SCZ vs BD genome-wide association study (GWAS) 134 significantly correlated with SCZ or BD diagnosis in an independent sample (Ruderfer et al., 2014), 135 providing the first evidence that differences between the disorders also have a genetic basis. An enrichment of rare, moderate to highly penetrant copy number variants (CNVs) and de novo CNVs 136 137 are seen in SCZ patients(CNV and Schizophrenia Working Groups of the Psychiatric Genomics 138 Consortium, 2017; Gulsuner and McClellan, 2015; Kirov et al., 2012; Stone et al., 2008; 139 Szatkiewicz et al., 2014), while, the involvement of CNVs in BD is less clear(Green et al., 2016). 140 Although the role of *de novo* single nucleotide variants in BD and SCZ has been investigated in 141 only a handful of studies, enrichment in pathways associated with the postsynaptic density has 142 been reported for SCZ, but not BD(Fromer et al., 2014; Kataoka et al., 2016). Identifying disorder-143 specific variants and quantifying the contribution of genetic variation to specific symptom 144 dimensions remain important open questions. Characterizing these genetic differences will 145 facilitate an understanding of the dimensions of the disorders instead of the dichotomous diagnosis.

For example, we have shown that SCZ patients with greater manic symptoms have higher polygenic risk for BD(Ruderfer et al., 2014). These findings demonstrate shared genetic underpinnings for symptoms across disorders and may enable us to characterize patients by genetic liability to symptom dimensions thereby informing disease course and treatment.

Here, we utilize large collections of genotyped samples for BD and SCZ along with clinicallyrelevant measures identifying 28 subphenotypes to address three questions: 1) Are there specific variants, genes or pathways that are either shared by, or differentiate BD and SCZ? 2) Are the shared symptoms between these disorders driven by the same underlying genetic profiles? and 3) Can we demonstrate independent genetic signatures for subphenotypes within these disorders?

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#### 156 **Results**

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#### 158 Shared genetic contribution to BD and SCZ

159 We performed association analysis of BD and SCZ combined into a single phenotype, totaling 160 53,555 cases (20,129 BD, 33,426 SCZ) and 54,065 controls on 15.5 million SNP allele dosages 161 imputed from 1000 genomes phase 3(The 1000 Genomes Project Consortium, 2015). Logistic 162 regression was performed controlling for 13 principal components of ancestry, study sites and 163 genotyping platform. We identified 11,231 SNPs with p-value below our genome-wide significance (GWS) threshold of  $5x10^{-8}$ . After grouping SNPs in linkage disequilibrium with each 164 165 other  $(r^2 > 0.2)$ , 114 genomic risk loci remained. For the most significant variant in each of the 114 166 GWS loci, we performed conditional analysis with any GWS hit within 1Mb of the extent of the 167 locus from the previously performed single disease GWAS of SCZ(Schizophrenia Working Group of the Psychiatric Genomics Consortium, 2014) and BD(Stahl et al., 2017) and identified 32 loci 168

169 that were independently significant defined strictly as no single disease locus within 1Mb or a 170 GWS p-value after conditional analysis (Supplementary Table 1). We further performed gene-set 171 based tests using MAGMA(Leeuw et al., 2015) across 10,891 curated pathways(Watanabe et al., 2017) and identified 8 pathways surpassing Bonferroni correction ( $p < 4.6 \times 10^{-6}$ ) with all but one 172 173 pathway implicating synaptic and neuronal biology (Supplementary Table 2a). Establishing 174 independent controls (see Methods) allowed us to perform disorder-specific GWAS in 20,129 BD 175 cases vs 21,524 BD controls and 33,426 SCZ cases and 32,541 SCZ controls. Using these results, 176 we compared effect sizes of these 114 loci across each disorder independently showing the subsets 177 of variants that had larger effects in SCZ compared to BD and vice versa (Figure 1a).

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#### 179 Differentiating genetic contribution to BD and SCZ

180 To identify loci with divergent effects on BD and SCZ, we performed an association analysis 181 comparing 23,585 SCZ cases with 15,270 BD cases matched for shared ancestry and genotyping 182 platform (see Methods, Figure 1b, Table 1). Two genome-wide significant loci were identified, the 183 most significant of which was rs56355601 located on chromosome 1 at position 173,811,455 184 within an intron of DARS2 (Supplementary Figure 1). The second most significant locus was 185 rs200005157, a four base-pair insertion/deletion, on chromosome 20 at position 47638976 in an 186 intron of ARFGEF2 (Supplementary Figure 2). For both variants, the minor allele frequency was 187 higher in BD cases than SCZ cases and disease-specific GWAS showed opposite directions of 188 effect when compared to controls. We sought to identify additional disease-specific loci by 189 comprehensively incorporating expression information with association results to perform fine-190 mapping and identify novel variants(Gamazon et al., 2015; Giambartolomei et al., 2014; Gusev et 191 al., 2016; He et al., 2013). Here, we applied the summary-data-based Mendelian randomization

192 (SMR) method(Zhu et al., 2016) (see Methods) utilizing the cis-QTLs derived from peripheral 193 blood(Westra et al., 2013), human dorsolateral prefrontal cortex (DLPFC)(Fromer et al., 2016) 194 from the Common Mind Consortium and 11 brain regions from the GTEx consortium(Consortium, 195 2015). We identified one SNP-probe combination that surpassed the threshold for genome-wide 196 significance in blood but was also the most significant finding in brain. We found that SNP 197 rs4793172 in gene *DCAKD* is associated with SCZ vs BD analysis ( $p_{GWAS} = 2.8 \times 10^{-6}$ ) and is an eQTL for probe ILMN 1811648 ( $p_{eOTL} = 2.9 \times 10^{-168}$ ), resulting in  $p_{SMR} = 4.1 \times 10^{-6}$  in blood ( $p_{eOTL}$ ) 198 =  $2.9 \times 10^{-25}$ ,  $p_{SMR} = 2.0 \times 10^{-5}$  in DLFC, and  $p_{eQTL} = 4.6 \times 10^{-15}$ ,  $p_{SMR} = 6.0 \times 10^{-5}$  in GTEx cerebellar 199 200 hemisphere) (Supplementary Table 3, Supplementary Figure 3) and shows no evidence of 201 heterogeneity ( $p_{HET} = 0.66$ ) which implies only a single causal variant in the locus.

202 In an effort to prioritize genes for the two GWS loci from the GWAS, we performed fine-203 mapping(Benner et al., 2016) using an LD map derived from a majority of the control samples. 204 We then performed SMR on each of the variants with causal probability greater than 1% using all 205 eQTLs from the CommonMind Consortium DLPFC reference. All the most likely causal variants 206 were shown to most significantly regulate the same gene suggesting *CSE1L* is the most likely relevant gene on chromosome 20 (rs200005157: causal probability=0.21, p<sub>GWAS</sub>=2.4x10<sup>-8</sup>, p<sub>eOTL</sub> 207  $3x10^{-8}$ , p<sub>SMR</sub>=8.5x10<sup>-5</sup>, p<sub>HET</sub>=0.34). For the locus on chromosome 1, SLC9C2 is the most 208 209 significantly regulated gene. However, a highly significant heterogeneity test indicates a complex 210 genetic architecture making it difficult to infer a causal role for the associated SNP. Therefore, DARS2 presents as the most likely relevant gene on chromosome 1 (rs56355601:  $p_{GWAS}$ =5.6x10<sup>-9</sup>, 211 causal probability=0.079,  $p_{eQTL}$  7.4x10<sup>-13</sup>,  $p_{SMR}$ =6.17x10<sup>-6</sup>,  $p_{HET}$ =0.03). We note however, that in 212 213 both cases there are less associated variants that are stronger eQTLs for these genes complicating 214 a straightforward causal interpretation. Finally, using the same gene-set test used for the combined

- analysis GO biological process "response to potassium ion" ( $p=1.6x10^{-6}$ ) was the only pathway surpassing our Bonferroni corrected significance threshold (Supplementary Table 2b).
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#### 218 Regional joint association

219 We expanded our efforts to identify disorder-specific genomic regions by jointly analyzing 220 independent GWAS results from BD and SCZ(Pickrell et al., 2016). The genome was split into 221 1,703 previously defined approximately LD independent regions(Berisa and Pickrell, 2015). 222 Thirteen percent, or 223 regions, had a posterior probability greater than 0.5 of having a causal 223 variant for at least one disorder. Of these, 132 best fit the model of a shared causal variant 224 influencing both BD and SCZ, 88 were most likely specific to SCZ, 3 demonstrated evidence of 225 two independent variants (with one impacting each of the two disorders) and none were BD-226 specific. Of note, this approach calculates a prior probability that any given region is disease-227 specific and from these data the probability of having a BD specific region was 0.1% compared to 228 15% for SCZ, likely a result of increased power from the larger SCZ sample size and/or a 229 difference in genetic architecture between these disorders.

230 The 114 GWS SNPs from the combined BD and SCZ GWAS localized into 99 independent 231 regions (13 regions had multiple GWS SNPs), of which 78 (79%) were shared with a posterior 232 probability of greater than 0.5. Sixty regions had at least one GWS SNP in the independent SCZ 233 GWAS, of which 30 (50%) are shared and 8 regions contained a GWS SNP in the independent 234 BD GWAS, of which 6 (75%) are shared using the same definition. For the three regions showing 235 evidence for independent variants, two had highly non-overlapping association signals in the same 236 region stemming from independent variants. The third, on chromosome 19 presented a different 237 scenario where association signals were overlapping. The most significant variant in BD was

rs111444407 (chr19:19358207, p = 8.67x10<sup>-10</sup>) and for SCZ was rs2315283 (chr19:19480575, 238 239  $p=4.41 \times 10^{-7}$ ). After conditioning on the most significant variant in the other disorder, the 240 association signals of the most significant variant in BD and SCZ were largely unchanged (BD rs111444407 =  $1.3 \times 10^{-9}$ , SCZ rs2315283 p= $6.7 \times 10^{-5}$ ). We further calculated the probability of each 241 242 variant in the region being causal for both BD and SCZ(Benner et al., 2016) and found no 243 correlation (r= -0.00016). The most significant variants had the highest posterior probability of 244 being causal (SCZ: rs2315283, prob = 0.02, BD: rs111444407, prob = 0.16). Both variants most 245 significantly regulate the expression of GATAD2A in brain (Fromer et al., 2016) but in opposite directions (rs111444407  $p_{eOTL} = 6x10^{-15}$ , beta = 0.105; rs2315283  $p_{eOTL} = 1.5x10^{-28}$ , beta = -0.11). 246

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#### 248 **Regional SNP-heritability estimation**

Across the genome, regional SNP-heritabilities  $(h^2_{snp})$  were estimated separately for SCZ and 249 250 BD(Shi et al., 2016) and were found to be moderately correlated (r=0.25). We next defined risk 251 regions as those containing the most associated SNP for each GWS locus. In total, there were 101 252 SCZ risk regions from the 105 autosomal GWS loci reported previously(Schizophrenia Working 253 Group of the Psychiatric Genomics Consortium, 2014) and 29 BD risk regions from 30 GWS loci 254 reported previously(Stahl et al., 2017). Ten regions were risk regions for both BD and SCZ 255 comprising 33% of BD risk regions and 10% of SCZ risk regions. We further stratified regional  $h_{snp}^2$  by whether a region was a risk region in one disorder, none or both (Supplementary Figure 256 257 4). Since the discovery data for the regions overlapped with the data used for the heritability 258 estimation, we expected within-disorder analyses to show significant results. In risk regions 259 specific to SCZ (n=91) there was a significant increase in regional  $h_{snp}^2$  in SCZ, as expected (p = 1.1x10<sup>-22</sup>), but also in BD ( $p = 1.2x10^{-6}$ ). In risk regions specific to BD (n=19), significantly 260

increased regional  $h_{snp}^2$  was observed in BD, as expected (p = 0.0007), but not in SCZ (p = 0.89). 261 262 Risk regions shared by both disorders had significantly higher  $h_{snp}^2$  in both disorders, as expected 263 (BD p =  $5.3 \times 10^{-5}$ , SCZ p = 0.006), compared to non-risk regions. However, we observed a 264 significant increase in BD  $h_{snp}^2$  in shared risk regions compared to BD risk regions (BD p = 0.003) 265 but not SCZ  $h_{snp}^2$  for shared risk regions compared to SCZ risk regions (p = 0.62). Using a less stringent p-value threshold for defining risk regions ( $p < 5x10^{-6}$ ), thereby substantially increasing 266 267 the number of regions, resulted in similar results. Seven regions contributed to substantially higher 268  $h_{snp}^2$  in SCZ compared to BD but no region showed the inverse pattern. Of these regions, all but 269 one was in the major histocompatibility region (MHC), the sole novel region was chr10:104380410-106695047 with regional  $h_{snp}^2 = 0.0019$  in SCZ and  $h_{snp}^2 = 0.00063$  in BD. 270

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#### 272 **Polygenic dissection of subphenotypes**

273 Subphenotypes were collected for a subset of patients with either BD or SCZ (see Methods). For 274 SCZ, we had clinical quantitative measurements of manic, depressive, positive and negative 275 symptoms generated from factor analysis of multiple instruments as described previously(Ruderfer 276 et al., 2014) but in larger sample sizes (n=6908, 6907, 8259, 8355 respectively). For BD, 24 277 subphenotypes were collected among nearly 13,000 cases in distinct categories including 278 comorbidities, clinical information such as rapid cycling and psychotic features as well as 279 additional disease course data such as age of onset and number of hospitalizations. For each BD 280 or SCZ patient, we calculated a polygenic risk score (PRS) using all SNPs, from each of the four 281 main GWAS analyses (BD+SCZ, BD, SCZ and SCZvsBD). We then used regression analysis 282 including principal components and site to assess the relationship between each subphenotype and 283 the 4 PRS. Specifically, we tested whether polygenic risk scores of BD+SCZ, BD, SCZ or

SCZvsBD were correlated with each of these subphenotypes separately within BD and SCZ cases. When testing if the variance explained by the PRS was different from zero, we applied a significance cutoff of p < 0.0004 based on Bonferroni correction for 112 tests. In total, we identified 6 significant results after correction (Figure 2, Table 2).

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289 A significant positive correlation existed between BD PRS and manic symptoms in SCZ cases as seen previously(Ruderfer et al., 2014) ( $p=2x10^{-5}$ , t=4.26) and BD PRS and psychotic features in 290 291 BD patients ( $p=5.3 \times 10^{-5}$ , t=4.04). A significant increase in SCZ PRS was seen for BD cases with versus without psychotic features ( $p=1.2x10^{-10}$ , t=6.45) and patients with increased negative 292 293 symptoms in SCZ patients (p=3.60x10<sup>-6</sup>, t=4.64). The BD+SCZ vs controls PRS was significantly associated with psychotic features in BD ( $p=7.9 \times 10^{-13}$ , t=7.17) and negative symptoms in SCZ 294 295  $(p=1.5 \times 10^{-5}, t=4.33)$ . The next two most significant results which did not survive our conservative 296 correction were both indicative of a more severe course in BD: increased BD+SCZ PRS with increased numbers of hospitalizations in BD cases ( $p=4.2x10^{-4}$ , t=3.53) and increased SCZ PRS 297 with earlier onset of BD ( $p=7.9 \times 10^{-4}$ , t=-3.36). We assessed the role of BD subtype on the 298 299 correlation between SCZ PRS and psychotic features and identified a significant correlation when 300 restricted to only BD type I cases indicating the result was not likely driven by BD patients with a schizoaffective subtype (BDI: 3,763 with psychosis, 2,629 without, p=1.55x10<sup>-5</sup>, Supplementary 301 302 Table 4).

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We performed a GWAS for all 8 quantitative subphenotypes and 9 binary subphenotypes with at least 1,000 cases and calculated heritability and genetic correlation with BD and SCZ. Only two subphenotypes had significant  $h_{snp}^2$  estimates using LD-score regression(Bulik-Sullivan et al.,

2015) both in BD: psychotic features in BD ( $h_{snp}^2=0.15$ , SE=0.06) and suicide attempt ( $h_{snp}^2=0.25$ , 307 308 SE=0.1). Only psychotic features demonstrated a significant genetic correlation with SCZ 309  $(r_g=0.34, SE=0.13, p=0.009)$ . The significant genetic correlation demonstrates a genome-wide 310 relationship between common variants contributing to SCZ risk and those contributing to 311 psychotic features in BD cases. We tested whether the most significantly associated SCZ loci 312 contributed directly to psychotic features in BD. One hundred of the 105 autosomal genome-wide 313 significant SCZ SNPs previously published(Schizophrenia Working Group of the Psychiatric 314 Genomics Consortium, 2014) were in our dataset after QC and 60 were in the same direction of 315 effect for risk of psychotic features in BD (p=0.028, one-sided binomial-test).

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#### 317

#### 318 **Discussion**

319 Here we present a genetic dissection of bipolar disorder and schizophrenia from over 100,000 320 genotyped subjects. Consistent with earlier results(Cross-Disorder Group of the Psychiatric 321 Genomics Consortium, 2013), we found extensive genetic sharing between these two disorders, 322 identifying 114 genome-wide significant loci contributing to both disorders of which 32 are novel. 323 These findings point to the relevance of neuronal and synaptic biology for the shared genetic 324 substrate of these disorders. However, despite this degree of sharing, we identified several loci that 325 significantly differentiated between the two disorders, having opposite directions of effect. We 326 also found polygenic components that significantly correlated from one disorder to symptoms of 327 the other.

328

Two GWS loci were identified from the case only SCZ versus BD analysis providing opportunities
to inform the underlying biological distinctions between BD and SCZ. The most significant locus

331 implicates DARS2 (coding for the mitochondrial Aspartate-tRNA ligase) which is highly expressed 332 in the brain and significantly regulated by the most significant SNP rs56355601 ( $p_{eOTL}=2.5 \times 10^{-11}$ ). 333 Homozygous mutations in DARS2 are responsible for leukoencephalopathy with brainstem and 334 spinal cord involvement and lactate elevation (LBSL), which was characterized by neurological 335 symptoms such as psychomotor developmental delay, cerebellar ataxia and delayed mental 336 development(Yamashita et al., 2013, p. 2). Based on methylation analysis from the prefrontal 337 cortex of stress models (rats and monkeys) and from peripheral samples (in monkeys and human 338 newborns), DARS2, among others, has been suggested as a potential molecular marker of early-339 life stress and vulnerability to psychiatric disorders(Luoni et al., 2016). The second most 340 significant locus implicates CSE1L, a nuclear transport factor that plays a role in cellular 341 proliferation as well as in apoptosis(Bera et al., 2001). Intronic SNPs in CSE1L have been 342 associated with subjective well-being(Okbay et al., 2016) and, nominally to antidepressant 343 response(Li et al., 2016). More interestingly, CSE1L is a potential target gene of miR-137, one of 344 the well-known schizophrenia risk loci(Schizophrenia Working Group of the Psychiatric 345 Genomics Consortium, 2014), which is able to negatively regulate CSE1L by interacting with 346 complementary sequences in the 3' UTR of CSE1L(Li et al., 2013). Although falling short of 347 genome-wide significance, the third most significant locus implicates ARNTL (Aryl Hydrocarbon 348 Receptor Nuclear Translocator Like), which is a core component of the circadian clock. ARNTL 349 has been previously hypothesized for relevance in bipolar disorder, (Yang et al., 2008) although 350 human genetic evidence is currently limited(Byrne et al., 2014).

351

The ability to generate transcriptional data on multiple tissues across many individuals using RNAsequencing has provided detailed information on the role common variants play in regulating

expression of specific genes in specific tissues. These eQTLs can be integrated with the genetic 354 355 association data from GWAS to inform on the relationship between variant association and variant 356 regulation of expression for each gene. Performing this integration, we identified a third genome-357 wide significant finding in DCAKD. The gene codes for Dephospho-CoA Kinase Domain 358 Containing protein, a member of the human postsynaptic density proteome from human 359 neocortex(Bayés et al., 2011). In the mouse cortical synaptoproteome DCAKD is among the 360 proteins with the highest changes between juvenile postnatal days and adult stage, suggesting a 361 putative role in brain development(Gonzalez-Lozano et al., 2016; Moczulska et al., 2014). 362 Discerning between pleiotropy (variant independently regulates expression and alters risk to 363 disease) from causality (variant regulates expression which thereby alters risk to disease) through 364 statistical analysis alone is difficult, this analytical approach is stringent in excluding loci where 365 colocalised SNP-phenotype and SNP-expression associations may reflect confounding driven by 366 linkage disequilibrium (LD) (one variant regulates expression and a different variant alters risk but 367 the variants in the region are in LD). Hence, this approach utilizes currently available data to 368 prioritize genes, including direction of effect, for functional follow-up. These analyses will become 369 more powered with increased sample sizes for both phenotype and eQTL data sets.

370

Performing pathway analysis based on the full association results shows enrichment of genes involved in response to potassium ions, including potassium voltage-gated channel subfamily members and a number of genes regulated by cellular potassium concentration. This is in line with previous genetic evidence pointing to a key etiologic role of potassium channels, in particular, in BD(Judy and Zandi, 2013), which could be explained by their role in multiple neurobiological mechanisms involved in the development of psychiatric disorders such as regulation of thedopaminergic circuits, synaptic plasticity, and myelination(Balaraman et al., 2015).

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379 We further assessed the contribution of regions of the genome to each disorder through joint 380 regional association and heritability estimation. These results point to an additional locus that may 381 contribute differentially to liability to BD and SCZ. The region on chr19 shows overlapping 382 association peaks that are driven by independent causal variants for each disorder. Both variants 383 significantly regulate the same gene GATAD2A but in opposite directions. GATAD2A is a 384 transcriptional repressor, which is targeted by MBD2 and is involved in methylation-dependent 385 gene silencing. The protein is part of the large NuRD (nucleosome remodeling and deacetylase) 386 complex, for which also HDAC1/2 are essential components. NurD complex proteins have been 387 associated with autism(Li et al., 2015). Their members, including GATAD2A, display preferential 388 expression in fetal brain development(Li et al., 2015) and in recent work has been implicated in 389 SCZ through open chromatin(Fullard et al., n.d.). Further, p66α (mouse GATAD2A) was recently 390 shown to participate in memory preservation through long-lasting histone modification in 391 hippocampal memory-activated neurons(Ding et al., 2017). SNP-heritability appears to be 392 consistently shared across regions and chromosomes between these two disorders. Regions with 393 GWS loci often explain higher proportions of heritability as expected. When looking at the effect 394 on heritability of the presence of a GWS locus in the other disorder, we identified a significant 395 increase in BD heritability for regions containing a GWS locus for SCZ but no significant increase 396 in SCZ heritability in regions having a BD one. This result suggests a directionality to the genetic 397 sharing of these disorders with a larger proportion of BD loci being specific to BD. However, we 398 cannot exclude that the asymmetry of results may reflect less power of discovery for BD than SCZ.

399 The degree to which power and subphenotypes contribute to this result requires further400 examination.

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402 We note that as with nearly all GWAS findings, the calculated population-based effect sizes of the 403 variants identified here are small and independently explain only a modest fraction to the 404 heritability of these disorders. The identification of these variants is dependent on the ability to 405 have highly accurate allele frequency estimates that can only be ascertained from large sample 406 sizes. As sample sizes get larger the power to identify variants of smaller effect increases meaning 407 that increasing sample size results in the identification of variants of smaller effect. However, a 408 small population effect size does not exclude the possibility of a substantially larger effect on 409 molecular phenotypes nor does it preclude the utility of association regions in understanding 410 biology or having a clinical impact. Efforts following up GWAS results to date have demonstrated 411 the value of these findings in pointing to genes that can aid in understanding the underlying biology 412 of the trait(Claussnitzer et al., 2015; Mohanan et al., 2018; Sekar et al., 2016). Further, there is a 413 clear relationship between GWAS results of a phenotype and gene targets of drugs that treat that 414 phenotype pointing to the potential for improved therapeutic understanding(Nelson et al., 2015; 415 Ruderfer et al., 2016). A major challenge of GWAS is the sheer number of findings and the 416 substantial time/cost required for functional follow up of these findings in the classical paradigms 417 used for genes causal for monogenic disorders. In silico bioinformatic analyses (such as SMR used 418 here) that integrate GWAS results with 'omics data (transcription, protein, epigenetic, etc.) have 419 the potential to put a clearer biological focus on GWAS results. Such analyses can become more 420 complex as more reference omics data sets (with genome-wide genotyping) become available. 421 Additional analytical efforts will be required to facilitate the transition from GWAS to biology but substantial data has shown there is much to be learned from these variants despite their smalleffects(Visscher et al., 2017).

424

425 We have now identified multiple genomic signatures that correlate between one disorder and a 426 clinical symptom in the other disorder, illustrating genetic components underlying particular 427 symptom dimensions within these disorders. Medical symptoms, including those seen in 428 psychiatric disorders, can manifest through a multitude of causes. The classic example often used 429 is headache for which many different paths lead to the same symptom. Psychiatric symptoms also 430 have many potential causes. For example, symptoms of psychosis can be the result of highly 431 heritable diseases such as BD and SCZ but also infectious and neurodegenerative diseases, 432 sleep/sensory deprivation or psychedelic drugs. Demonstrating a shared biological underpinning 433 to these symptoms suggests they could be treated through modulating the same pathway. As 434 previously shown, we find a significant positive correlation between the PRS of BD and manic 435 symptoms in SCZ. We also demonstrate that BD cases with psychotic features carry a significantly 436 higher SCZ PRS than BD cases without psychotic features and this result is not driven by the 437 schizoaffective BD subtype. Further, we show that increased PRS is associated with more severe 438 illness. This is true for BD with psychotic features having increased SCZ PRS, earlier onset BD 439 having higher SCZ PRS and cases with higher BD+SCZ PRS having a larger number of 440 hospitalizations. We demonstrated that psychotic features within BD is a heritable trait and GWS 441 loci for SCZ have a consistent direction of effect in psychotic features in BD, demonstrating the 442 potential to study psychosis more directly to identify variants contributing to that symptom 443 dimension.

444

445 This work illustrates the utility of genetic data, in aggregate, at dissecting symptom heterogeneity 446 among related disorders and suggests that further work could aid in characterizing patients for 447 more personalized treatment. Genetic risk scores have demonstrated their ability to inform and 448 predict pathology (Cleynen et al., 2016) and more recently have been shown to be able to identify 449 patients with risk equivalent to monogenic variants (Khera et al., 2017). In psychiatry, we lack 450 objective biological measurements (biomarkers) with which to assess the ability of a genetic 451 signature to predict or inform. Lacking diagnostic pathology for psychiatric disorders leaves a 452 genuine opportunity for the genetics to drive diagnosis and treatment to a much larger degree than 453 in other domains. One potential model assumes that each individual has a quantitative loading of 454 a series of symptom dimensions (i.e. manic, psychotic, cognitive, etc.) and that these symptoms 455 can be assessed at the genetic level to characterize a patient's dysfunction and used to inform 456 disease course and optimal treatment. Making this a reality will require more detailed information 457 on disease course and outcomes. For example, if treatment response data existed for these samples 458 one could ask whether a genetic loading for psychosis was correlated with response to treatment. 459 Initial work has already shown the potential of this approach using a SCZ PRS to inform lithium 460 response in BD(Amare et al., 2018). Ultimately, the goal will be to quantify multiple genetic 461 loadings of each individual's illness and use those measures to inform treatment based on the 462 outcomes of previous individuals with similar profiles.

463

In conclusion, we present a detailed genetic dissection of BD and SCZ pointing to substantial shared genetic risk but also demonstrating that specific loci contribute to the phenotypic differences of these disorders. We show that genetic risk scores can correspond to symptoms within and across disorders. Finally, we present data that points to these disorders being neither

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independent nor the same but sharing particular symptom dimensions that can be captured fromthe genetics and used to characterize patients to ultimately inform diagnosis and treatment.

470

#### 471 Author Contributions:

472 DMR, PS and KSK managed and organized the group. DMR, SR, JB, EAS, JMWP, NM, AWC,

473 APSO, LMOL and VT contributed to analyses. Subphenotype collection and organization was led

474 by AM and AHF. Initial manuscript was drafted by DMR, ED, ADF, SP, JLK. Manuscript

475 contributions and interpretation of results was provided by DMR, ED, SHL, MCO, PFS, RAO,

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- 477 processing for the contributing components of the study. All other authors saw, had the opportunity

478 to comment on, and approved the final draft.

480

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- 578 The authors declare no competing interests.
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### 899 Figure Legends

900

### 901 Figure 1. Associated Genomic Loci Shared and Divergent Between BD and SCZ

- a) Odds ratios (OR) from independent data sets of BD (blue) and SCZ (red) for each of the 114
- 903 genome-wide significant variants in the BD and SCZ vs controls GWAS. b) Manhattan plot for

904 SCZ vs BD GWAS.

905

#### 906 Figure 2. Polygenic Risk Score Dissection of Clinical Symptom Dimensions

907 Effect size (calculated by dividing regression estimate by standard error) from regression analysis 908 including ancestry covariates for each subphenotype and PRS for BD (x-axis) and SCZ (y-axis). 909 Point size represents -log10(p-value) with SCZ (red) and BD (blue). Numbered subphenotypes 910 are 1) comorbid migraine, 2) panic attacks 3) suicide attempt 4) mixed states 5) rapid cycling 6) 911 comorbid eating disorder 7) comorbid OCD 8) year of birth 9) suicide ideation 10) panic disorder 912 11) number of suicide attempts 12) depressive symptoms (SCZ) 13) episodes depressive 14) 913 episodes total 15) positive symptoms (SCZ) 16) irritable mania 17) age of onset depression 18) 914 family history 19) episodes mixed mania 20) unipolar mania 21) alcohol substance dependence 915 22) age of onset mania 23) age at interview 24) number of hospitalizations. All subphenotypes are 916 in BD except those labeled (SCZ).

917

#### 918 Table Legends

919

#### 920 Table 1. Most Significant Associated Loci from SCZ vs BD GWAS

921 Association results for the five most significant variants in the SCZ vs BD GWAS with the top 922 two being genome-wide significant. Each variant includes results from the independent BD vs 923 controls and SCZ vs controls GWAS and the comparable p-value from a heterogeneity test when 924 performing a two cohort meta-analysis of SCZ and BD.

925

#### 926 Table 2. Complete Results of Polygenic Risk Score Dissection Analysis

927	Polygenic scoring results of all four GWAS phenotypes (BD+SCZ vs controls, BD vs controls,
928	SCZ vs controls and SCZ vs BD) and 24 subphenotypes from BD and 4 subphenotypes from SCZ,
929	rows without case/control counts are quantitative measures. Significance and effects are from
930	regression analysis of subphenotype on PRS including principal components of ancestry and site
931	as covariates. Effect is the regression estimate divided by the standard error.
932	
933	Supplementary Figure Legends
934	
935	Figure S1. Related to Figure 1b. Regional Association Plot and Forest Plot for the First
936	Genome-wide Significant Hit in the SCZ vs BD GWAS.
937	Figure S2. Related to Figure 1b. Regional Association Plot and Forest Plot for the Second
938	Genome-wide Significant Hit in the SCZ vs BD GWAS.
939	
940	
941	Figure S3. Related to Summary-data-based Mendelian Randomization. Detailed Association
942	of DCAKD from SMR.
943	Results at the DCAKD locus from SMR analysis of SCZ vs BD. Top plot, brown dots represent
944	the P values for SNPs from SCZ vs BD GWAS, diamonds represent the P values for probes from
945	the SMR test. Bottom plot, the $eQTL P$ values of SNPs from the Westra study for the
946	ILMN_1811648 probe tagging DCAKD. The top and bottom plots include all the SNPs available
947	in the region in the GWAS and eQTL summary data, respectively, rather than only the SNPs
948	common to both data sets. Highlighted in red is the gene (DCAKD) that passed the SMR and

HEIDI tests.

950

#### 951 Figure S4. Related to Regional SNP-heritability estimation. Heritability Estimates for BD

#### 952 and SCZ in Genome-wide Significant Regions of BD and SCZ.

- 953 Regional SNP-heritability estimates for SCZ and BD stratified by whether the region contains the
- 954 most significant variant in a genome-wide significant locus in BD, SCZ, neither or both.

955

956

957 STAR Methods

#### 958 CONTACT FOR REAGENT AND RESOURCE SHARING

959 Genotype and phenotype data use is restricted and governed by the Psychiatric Genetics 960 Consortium. Further information and requests for analytical results or additional information 961 should be directed to and will be fulfilled by the Lead Contact, Douglas Ruderfer 962 (douglas.ruderfer@vanderbilt.edu).

963

#### 964 SUBJECT DETAILS

#### 965 Genotyped Sample Description

966 SCZ samples are a substantial subset of those analyzed previously(Schizophrenia Working Group

967 of the Psychiatric Genomics Consortium, 2014). BD samples are the newest collection from

968 Psychiatric Genomics Consortium Bipolar Disorder Working Group(Stahl et al., 2017).

969 Below we provide information on the individual samples used here as provided by the original

- 970 PGC disorder publications. Additionally, most studies have been described in detail in the citations
- 971 provided. The boldfaced first line for each sample is study PI, PubMed ID, country (study name),
- 972 and the PGC internal tag or study identifier.

973

#### 974 European ancestry, case-control design

- 975 Schizophrenia
- 976 Adolfsson, R | NP | Umeå, Sweden | scz\_umeb\_eur

#### 977 Adolfsson, R | NP | Umeå, Sweden | scz\_umes\_eur

978 Cases of European ancestry were ascertained from multiple different studies of schizophrenia 979 (1992-2009). The diagnostic processes were similar between studies, and the final diagnosis is a 980 best-estimate consensus lifetime diagnosis based on multiple sources of information such as 981 clinical evaluation by research psychiatrists, different types of semi-structured interviews made by 982 trained research nurses and research psychiatrists, medical records, course of the disease and data 983 from multiple informants. Diagnosis was made in accordance with the Diagnostic and Statistical 984 Manual of Mental Disorders-Version IV (DSM-IV) or International Classification of Diseases, 985 10th Revision (ICD-10) criteria. Controls were recruited from the Betula study, an ongoing 986 longitudinal, prospective, population-based study from the same geographic area (North Sweden) 987 that is studying aging, health, and cognition in adults. All subjects (cases and controls) participated 988 after giving written informed consent and the regional Ethical Review Board at the University of 989 Umeå approved all original studies and participation in the PGC. GWAS genotyping was 990 performed at Broad Institute.

#### 991 Andreassen, O | 19571808 | Norway (TOP) | scz\_top8\_eur

In the TOP study (Tematisk omrade psykoser), cases of European ancestry, born in Norway, were recruited from psychiatric hospitals in the Oslo region. Patients were diagnosed according to SCID and further ascertainment details have been reported. Healthy control subjects were randomly selected from statistical records of persons from the same catchment area as the patient groups. 996 All participants provided written informed consent and the human subjects protocol was approved

997 by the Norwegian Scientific-Ethical Committee and the Norwegian Data Protection Agency.

#### 998 Blackwood, D | 19571811 | Edinburgh, UK | scz\_edin\_eur

999 Cases and controls were recruited from the southeast of Scotland, and ascertainment has been 1000 previously described as part of the International Schizophrenia Consortium studies. All 1001 participating subjects gave written, informed consent and the human subjects protocol was 1002 approved by the Scotland A Research Ethics Committee. DNA samples were genotyped at the 1003 Broad Institute.

#### 1004 Børglum, A | 19571808 | Denmark | scz\_aarh\_eur

1005 DNA samples for all subjects were collected from blood spots systematically collected by the 1006 Danish Newborn Screening Biobank), with case/control status established using the Danish 1007 Psychiatric Central Register. Cases were diagnosed clinically according to ICD-10 criteria. 1008 Controls were selected to match the cases by birth cohort. The Danish Data Protection Agency and 1009 the ethics committees in Denmark approved the human subjects protocol.

#### 1010 Bramon | 23871474 | Seven countries (PEIC, WTCCC2) | scz\_pewb\_eur

#### 1011 Bramon | 23871474 | Spain (PEIC, WTCCC2) | scz\_pewb\_eur

1012 The Psychosis Endophenotypes International Consortium (PEIC) was part of WTCCC2. Samples 1013 were collected through seven centers in Europe and Australia (the Institute of Psychiatry, King's 1014 College London, London; GROUP (consisting of the University of Amsterdam, Amsterdam; the 1015 University of Groningen, Groningen; Maastricht University Medical Centre, Maastricht; and the 1016 University of Utrecht, Utrecht); the University of Western Australia, Perth; the Universidad de 1017 Cantabria, Santander; the University of Edinburgh, Edinburgh; Heidelberg University, Heidelberg 1018 and Ludwig-Maximilians-Universität München, Munich). To allow for a DSM-IV diagnosis to be 1019 ascertained or ruled out, all participants (including controls and unaffected family members) 1020 underwent a structured clinical interview with the Schedule for Affective Disorders and 1021 Schizophrenia (SADS), the Structured Clinical Interview for DSM Disorders (SCID), or the 1022 Schedules for Clinical Assessment in Neuropsychiatry (SCAN). We included cases with 1023 schizophrenia and schizoaffective disorder. Participants in all groups were excluded if they had a 1024 history of neurological disease or head injury resulting in loss of consciousness.

#### 1025 Buxbaum, J | 20489179 | New York, US & Israel | scz\_msaf\_eur

Samples contributed by Mount Sinai were derived from three cohorts. In all cohorts, ethical approval was obtained from all participating sites, and all subjects provided informed consent. Two of the cohorts were in a prior paper on copy number variation. One of the cohorts was from the Mount Sinai brain bank, where DNA was extracted from postmortem samples, and another comprised of patients ascertained in Israel. The third cohort included subjects more recently recruited through the Mount Sinai Conte Center.

#### 1032 Corvin, A | 19571811 | Ireland | scz\_dubl\_eur

1033 The case sample was collected primarily in the Dublin area and the ascertainment procedure has 1034 been previously described. The controls were recruited, from the same region through the Irish 1035 Blood Transfusion Services. All participants gave written, informed consent and the collections 1036 were approved through the Federated Dublin Hospitals and Irish Blood Transfusion Services 1037 Research Ethics Committees, respectively. DNA samples were genotyped at the Broad Institute.

#### 1038 Corvin, A; Riley, B | 22883433 | Ireland (WTCCC2) | scz\_irwt\_eur

1039 The case sample was recruited from the Republic of Ireland and Northern Ireland. All cases had 1040 four Irish grandparents and ascertainment details have been reported elsewhere. Ethics approval 1041 was obtained from all participating hospitals and centers. Controls were blood donors from the 1042 Irish Blood Transfusion Service, whose Ethics Committee approved the human subjects protocol.1043 All participants gave written informed consent. Samples were genotyped at Affymetrix (Santa

1044 Clara, California, US) laboratory as part of the WTCCC2 genotyping pipeline.

#### 1045 Ehrenreich, H | 20819981 | Germany (GRAS) | scz\_gras

The Gottingen Research Association for Schizophrenia (GRAS) collection included cases recruited across 23 German hospitals. Controls were unscreened blood donors recruited at the Georg-August-University according to national blood donation guidelines. Cases completed a structured clinical interview and were diagnosed with DSM-IV schizophrenia or schizoaffective disorder. The study was approved by the Georg-August-University ethics committee and local internal review boards of the participating centers. All participants gave written informed consent.

#### 1052 Esko, T | 15133739 | Estonia (EGCUT) | scz\_egcu\_eur

1053 The Estonian cohort comes from the population-based biobank of the Estonian Genome Project of 1054 University of Tartu (EGCUT). The project was conducted according to the Estonian Gene 1055 Research Act and all participants provided informed consent (www.biobank.ee). In total, 52,000 1056 individuals aged 18 years or older participated in this cohort (33% men, 67% women). The 1057 population distributions of the cohort reflect those of the Estonian population (83% Estonians, 1058 14% Russians and 3% other). General practitioners (GP) and physicians in the hospitals randomly 1059 recruited the participants. A Computer-Assisted Personal interview was conducted over 1-2 ours 1060 at doctors' offices. Data on demographics, genealogy, educational and occupational history, 1061 lifestyle and anthropometric and physiological data were assessed. Schizophrenia was diagnosed 1062 prior to the recruitment by a psychiatrist according to ICD-10 criteria and identified from the 1063 Estonian Biobank phenotype database. Controls were drawn from a larger pool of genotyped

- 1064 biobank samples by matching on gender, age and genetic ancestry. All the controls were 1065 population-based and have not been sampled for any specific disease.
- 1066 Esko, T; Li, Q; Dominici E | 15133739, 24166486 | J&J and Roche cases, EGCUT controls |
   1067 scz\_jr3a\_eur
- 1068 Esko, T; Li, Q; Domenici E | 15133739, 24166486 | J&J and Roche cases, EGCUT controls |
   1069 scz\_jr3b\_eur
- 1070 Esko, T; Li, Q; Domenici E | 15133739, 24166486 | J&J and Roche cases, EGCUT controls |
   1071 scz\_jri6\_eur
- 1072 Esko, T; Li, Q; Dominici E | 15133739, 24166486 | J&J and Roche cases cases, EGCUT
  1073 controls | scz\_jrsa\_eur
- 1074 Cases were collected by Johnson and Johnson (J&J) and Roche as part of clinical collaborations 1075 with hospitals and outpatient centers. Cases were diagnosed according to DSMIV criteria, with 1076 medical record review by a trained psychiatrist. There were reliability trials across centers for the 1077 J&J studies. The J& J cases were mostly collected in Eastern Europe, with most coming from 1078 Estonian and Russia (>100); intermediate numbers from Austria, the Czech Republic, Latvia, 1079 Lithuania, and Spain (50-100); and smaller collections from Bulgaria, Hungary, and Poland (<50). 1080 The Roche cases were assessed with a structured psychiatric assessment by trained interviewers. 1081 Most of the Eastern European controls were from the Estonian Biobank project (EGCUT) and 1082 were ancestrally matched with cases from the J&J sample.
- 1083 Gejman, P | 19571809 | US, Australia (MGS) | scz\_mgs2\_eur

European ancestry case samples were collected by the Molecular Genetics of Schizophrenia (MGS) collaboration across multiple sites in the USA and Australia as described in detail elsewhere. Cases gave written informed consent, and IRBs at each collecting site approved the human subjects protocol. A survey company (Knowledge Networks, under MGS guidance)
collected the European ancestry control sample and ascertainment is described in detail elsewhere.

1089 DNA samples were genotyped at the Broad Institute.

1090 Gurling, H | 19571811 | London, UK | scz\_uclo\_eur

All cases and controls were collected by University College London and had both parents from England, Scotland or Wales. All participants gave written informed consent and the U.K. National Health Service multicenter and local research ethics committee approved the human subjects protocol. Further details on ascertainment are available elsewhere. The samples were genotyped at the Broad Institute.

#### 1096 Jönsson, E | 19571808 | Sweden (Hubin) | scz\_ersw\_eur

1097 Cases were recruited from northwestern Stockholm County and ascertainment has been described 1098 previously. Cases gave informed consent and the human subjects protocol was approved by the 1099 ethical committees of the Karolinska Hospital and the Stockholm Regional Ethical Committee. 1100 Controls were recruited either among subjects previously participating in biological research at the 1101 Karolinska Institute or drawn from a representative register of the population of Stockholm 1102 County. All participants provided informed consent.

#### 1103 Kirov, G | Not published | Bulgaria | scz\_buls\_eur

All cases were recruited from Bulgaria and had a history of hospitalization for treatment of schizophrenia. Controls were recruited from the two largest cities in Bulgaria as previously described. All participants gave written informed consent and the study was approved by local ethics committees at the participating centers.

1108 Knight, J; Collier DA; Nisenbaum L| Not published | Canada (Toronto) -US(Lilly)-US

1109 (MIGen) scz\_lktu\_eur

37

Toronto cases were recruited by referral and advertisement. Diagnoses were made according to DSM-III or DSM-IV criteria following interview and medical record review. US cases were recruited from schizophrenia clinical trials in a range of settings as part of a trial with Eli Lilly. Diagnoses were made according to DSM-III or DSM-IV criteria following interview by psychiatrist and medical record review. No controls were sampled as part of the study, and ancestrally-matched controls were chosen from the Myocardial Infarction Genetics Consortium (MIGen, dbGaP ID phs000294.v1.p1) that was genotyped with the same SNP array.

#### 1117 Lencz, T; Darvasi A | 23325106 | Israel | scz\_ajsz\_eur

1118 Cases and controls were sampled from an Ashkenazi Jewish repository (Hebrew University 1119 Genetic Resource, http://hugr.huji.ac.il). Patients were recruited from hospitalized inpatients at 7 1120 medical centers in Israel and were diagnosed with DSM-IV schizophrenia or schizoaffective 1121 disorder. Controls were sampled through the Israeli Blood Bank and did not report any chronic 1122 disease or regularly prescribed medication at the time of assessment. Full ascertainment details 1123 have previously been reported. Local ethics committees and the National Genetic Committee of 1124 the Israeli Ministry of Health approved the studies and all participants gave informed, written 1125 consent.

#### 1126 Levinson, D | 22885689 | Six countries, WTCCC controls | scz\_lacw\_eur

1127 Cases collected as part of a larger pedigree-based study were partitioned into two subsamples. 1128 Cases with two genotyped parents were analyzed as trios (see PI Levinson, ms.scz\_lemu\_eur in 1129 the Trio section below). Unrelated cases who could not be used as part of a trio were included as 1130 a separate case-control analysis, using independent controls, matched by ancestry and genotyping 1131 array, from the Wellcome Trust Case Control Consortium. Cases were identified from different 1132 clinical settings (e.g. inpatients, outpatients and community facilities) in six countries (Australia, France, Germany, Ireland, UK, and the US). Diagnoses were established using semi-structured interviews, psychiatric records and informant reports. Case subjects were diagnosed with schizophrenia or schizoaffective disorder according to DSM-III-R criteria. All protocols were approved by loci IRBs, and all cases provided written informed consent.

#### 1137 Malhotra, A | 17522711 | New York, US | scz\_zhh1\_eur

1138 The case and control subjects were recruited in the New York metropolitan area and ascertainment 1139 methods have been described previously. All participants gave written, informed consent and the 1140 IRB of the North Shore-Long Island Jewish Health System approved the human subjects protocols.

1141 DNA was genotyped at Zucker Hillside.

#### 1142 Mowry, B | 21034186 | Australia | scz\_asrb\_eur

1143 These subjects were part of the Australian Schizophrenia Research Bank. The case sample was 1144 recruited in four Australian States (New South Wales, Queensland, Western Australia and 1145 Victoria) through hospital inpatient units, community mental health services, outpatient clinics and 1146 rehabilitation services, non-government mental illness support organizations, and, in the initial 1147 stages, through a large-scale, national, multi-media advertising campaign. This sample is 1148 comprised of 509 cases from larger metropolitan centers of Brisbane, Newcastle, Sydney, 1149 Melbourne, and Perth. Cases gave written informed consent, and the human subjects protocol was 1150 initially approved by the Hunter New England Area Health Research Committee and subsequently 1151 approved by relevant Institutional Ethics Committees in Brisbane, Sydney, Melbourne and Perth. 1152 Healthy controls were recruited through multi-media advertisements, and other sources. Controls 1153 were from the metropolitan centers of Brisbane, Newcastle, Sydney, Melbourne, and Perth. 1154 Controls gave written informed consent, and the human subjects protocol was approved by the 1155 Hunter New England Area Health Research Committee and Institutional Ethics Committees in 1156 Brisbane, Sydney, Melbourne and Perth. The samples were genotyped in two stages at the Hunter

1157 Medical Research Institute, University of Newcastle, Newcastle, Australia.

#### 1158 O'Donovan, M: Owen, M | 19571811 | Cardiff, UK | scz\_caws\_eur

The case sample included European ancestry schizophrenia cases recruited in the British Isles and described previously. All cases gave written informed consent to. The study was approved by the Multicentre Research Ethics Committee in Wales and Local Research Ethics Committees from all participating sites. The control sample used the Wellcome Trust CaseControl Consortium (WTCCC) sample described elsewhere, but included similar numbers of individuals from the 1958 British Birth Cohort and a panel of consenting blood donors (UK Blood Service). Samples were genotyped at Affymetrix service lab (San Francisco, USA).

#### 1166 O'Donovan, M: Owen, M: Walters, J | 22614287 | UK (CLOZUK) | scz\_clm2\_eur

#### 1167 O'Donovan, M: Owen, M: Walters, J | 22614287 | UK (CLOZUK) | scz\_clo3\_eur

1168 CLOZUK cases were taking the antipsychotic clozapine and had received a clinical diagnosis of 1169 treatment-resistant schizophrenia. Patients taking clozapine provide blood samples to allow 1170 detection of adverse drug-effects. Through collaboration with Novartis (the manufacturer of a 1171 proprietary form of clozapine, Clozaril), we acquired blood from people with treatment-resistant 1172 schizophrenia according to the clozapine registration forms completed by treating psychiatrists as 1173 previously reported. The samples were genotyped at the Broad Institute. The UK Multicentre 1174 Research Ethics Committee (MREC) approved the study. The controls were drawn from the 1175 WTCCC2 control samples (~3,000 from the 1958 British Birth Cohort and ~3,000 samples from 1176 the UK Blood Service Control Group). An additional 900 controls, held by Cardiff University, 1177 were recruited from the UK National Blood Transfusion Service. They were not specifically

screened for psychiatric illness. All control samples were from participants who provided informedconsent.

#### 1180 Ophoff, R | 19571808 | Netherlands | scz\_ucla\_eur

The case sample consisted of inpatients and outpatients recruited through psychiatric hospitals and institutions throughout the Netherlands. Cases with DSM-IV schizophrenia were included in the analysis. Further details on ascertainment are provided elsewhere. Controls came from the University Medical Centre Utrecht and were volunteers with no psychiatric history. Ethical approval was provided by local ethics committees and all participants gave written informed consent.

1187 Palotie, A | 19571808 | Finland | scz\_fi3m\_eur

#### 1188 Palotie, A | Not published | Finnish | scz\_fii6\_eur

Finnish cases were drawn from a nationwide collection of families with schizophrenia spectrum disorders. The control sample was derived from the Finnish Health 2000 survey. All participants provided written informed consent and approval was obtained from the ethics committees at each location.

1193 Pato, C | 19571811 | Portugal | scz\_port\_eur

Cases and controls lived in Portugal, the Azorean and Madeiran islands, or were the direct (firstor second-generation) Portugese immigrant population in the US, as previously described. Controls were not biologically related to cases. All participants gave written informed consent and the IRB of SUNY Upstate Medical University approved the protocol. The samples were genotyped at the Broad Institute.

1199 Petryshen, T | 24424392| Boston, US (CIDAR) | scz\_cims\_eur

1200 Cases were recruited from inpatient and outpatient settings in the Boston area by clinician referral, 1201 through review of medical records, or through advertisements in local media. Cases were 1202 diagnosed with DSM-IV schizophrenia through a structured clinical interview (SCID) by trained 1203 interviewers with review of medical records and a best estimate diagnostic procedure including 1204 reliability trials across interviewers. A psychiatrist or a PhD-level mental health professional made 1205 the final diagnostic determination. Controls were ascertained through local advertisements from 1206 the same geographical area. Ethical approval was provided by local ethics committees and all 1207 participants gave written informed consent.

#### 1208 Rietschel/Rujescu/Nöthen | 19571808 | Bonn/Mannheim, Germany | scz\_boco\_eur

1209 These German samples were collected by separate groups within the MooDS Consortium in 1210 Mannheim, Bonn, Munich and Jena. For the PGC analyses, the samples were combined by chip 1211 and ancestry. In Bonn/Mannheim, cases were ascertained as previously described. Controls were 1212 drawn from three population-based epidemiological studies (PopGen), the Cooperative Health 1213 Research in the Region of Augsburg (KORA) study, and the Heinz Nixdorf Recall (HNR) study. 1214 All participants gave written informed consent and the local ethics committees approved the 1215 human subjects protocols. Additional controls were randomly selected from a Munich-based 1216 community sample and screened for the presence of anxiety and affective disorders using the 1217 Composite International Diagnostic Screener. Only individuals negative for the above mentioned disorders were included in the sample. 1218

#### 1219 Rujescu, D | 19571808 | Munich, Germany | scz\_munc\_eur

For the Munich sample, cases were ascertained from the Munich area of Germany, as described previously. The controls were unrelated volunteers randomly selected from the general population of Munich. All were screened to exclude a history of psychosis/central neurological disease either 1223 personally or in a first-degree relative. All participants gave written informed consent and the local

1224 ethics committees approved the human subjects protocols.

#### 1225 St Clair, D | 19571811 | Aberdeen, UK | scz\_aber\_eur

Ascertainment and inclusion/exclusion criteria for cases and controls have been previously described. All participating subjects were born in the UK (95% Scotland) and gave written informed consent. Both local and multiregional academic ethical committee approved the human subjects protocol. The samples were genotyped at the Broad Institute.

#### 1230 Sullivan, PF | 18347602 | US (CATIE) | scz\_cati\_eur

Cases were collected as part of the Clinical Antipsychotics Trials of Intervention Effectiveness (CATIE) project and ascertainment was previously described. Participants were recruited from multiple sites in the USA with informed written consent and approval from the IRBs at each CATIE site and the University of North Carolina (Chapel Hill). The control subjects were collected by MGS (described above) and gave online informed consent and were fully anonymized. There was no overlap with controls included in the MGS collaboration sample.

- 1237 Sullivan, PF; Sklar P; Hultman C | 23974872 | Sweden | scz\_swe1\_eur
- 1238 Sullivan, PF; Sklar P; Hultman C | 23974872 | Sweden | scz\_s234\_eur
- 1239 Sullivan, PF; Sklar P; Hultman C | 23974872 | Sweden | scz\_swe5\_eur
- 1240 Sullivan, PF; Sklar P; Hultman C | 23974872 | Sweden | scz\_swe6\_eur

Samples from the Swedish Schizophrenia Study were collected in a multi-year project and genotypes in six batches (sw1-6). All procedures were approved by ethical committees at the Karolinska Institutet and the University of North Carolina, and all subjects provided written informed consent (or legal guardian consent and subject assent). All samples were genotyped at the Broad Institute. Cases with schizophrenia were identified via the Swedish Hospital Discharge 1246 Register which captures all public and private inpatient hospitalizations. The register is complete 1247 from 1987 and is augmented by psychiatric data from 1973-1986. The register contains 1248 International Classification of Disease discharge diagnoses made by attending physicians for each 1249 hospitalization. Case inclusion criteria included  $\geq 2$  hospitalizations with a discharge diagnosis of 1250 schizophrenia, both parents born in Scandinavia and age  $\geq 18$  years. Case exclusion criteria 1251 included hospital register diagnosis of any medical or psychiatric disorder mitigating a confident 1252 diagnosis of schizophrenia as determined by expert review. The validity of this case definition of 1253 schizophrenia was strongly supported by clinical, epidemiological, genetic epidemiological and 1254 genetic evidence. Controls were selected at random from Swedish population registers, with the 1255 goal of obtaining an appropriate control group and avoiding 'super-normal' controls. Control 1256 inclusion criteria included never being hospitalized for schizophrenia or bipolar disorder (given 1257 evidence of genetic overlap with schizophrenia), both parents born in Scandinavia and age of  $\geq 18$ 1258 years.

#### 1259 Walters, J | 21850710 | Cardiff, UK (CogUK) | scz\_cou3\_eur

Cases were recruited from community mental health teams in Wales and England on the basis of a clinical diagnosis of schizophrenia or schizoaffective disorder (depressed sub-type) as described previously. 35 Diagnosis was confirmed following a SCAN interview and review of case notes followed by consensus diagnosis according to DSM-IV criteria. The samples were genotyped at the Broad Institute. The UK Multicentre Research Ethics Committee (MREC) approved the study and all participants provided valid informed consent.

- 1266 Weinberger, D | 11381111 | NIMH CBDB | scz\_lie2\_eur
- 1267 Weinberger, D | 11381111 | NIMH CBDB | scz\_lie5\_eur

Subjects were recruited from the Clinical Brain Disorders Branch of the NIMH 'Sibling Study' as previously described. In brief, cases and controls gave informed consent and only participants of European ancestry were included in the current analysis. Cases completed a structured clinical interview and were diagnosed with schizophrenia-spectrum disorders. Samples were genotyped at the NIMH.

#### 1273 Wendland/Schubert | Pfizer | Not Published | Multiple countries | scz\_pfla\_eur

1274 Pfizer contributed anonymized individual genotypes for cases from seven multi-center 1275 randomized, double-blind efficacy and safety clinical trials (A1281063, A1281134, A1281148, 1276 A245-102, NRA7500001, NRA7500002, NRA7500003, and NRA7500004) as well as a set of 1277 purchased samples (NRA9000099). Trial samples were collected for antipsychotic medications 1278 across outpatient and inpatient treatment settings. All participating cases had a diagnosis of 1279 schizophrenia and were assessed using a structural clinical interview by trained interviewers, with 1280 systematic procedures to quality-control diagnostic accuracy and reliability trials across 1281 participating sites in the United States and internationally. Purchased blood samples were obtained 1282 from PrecisionMed International by Pharmacia and Upjohn Corporation, and were collected from 1283 diagnosed subjects with schizophrenia and schizoaffective disorder. All studies were reviewed by 1284 both central and local institutional review boards, depending on the study site, before recruitment 1285 of subjects started. Protocol amendments were approved while the study was in progress and 1286 before the data were unblinded. The studies were conducted in conformity with the U.S. Food and 1287 Drug Administration Code of Federal Regulations (21CFR, Part 50) and the Declaration of 1288 Helsinki and its amendments, and were consistent with Good Clinical Practice and the applicable 1289 regulatory requirements. Participants provided written informed consent before enrollment. An 1290 optional blood sample was collected from clinical trial subjects for pharmacogenetic analysis to

investigate potential associations between genetic variant drug response and general characteristics of schizophrenia and related disorders. Sample collection was not required for participation in the original clinical trials. The controls (A9011027) were recruited in a multi-site, cross-sectional, non-treatment prospective trial to collect data, including DNA, from cognitive normal and free of psychiatric diseases elderly subjects in the US. Subjects were specifically recruited to match the gender, age, and ethnicity information from the LEADe and UCSD MCI studies. The study described here is within the scope of patient consent.

1298 Werge, T | 19571808 | Denmark | scz\_denm\_eur

Cases were ascertained through psychiatric departments and twin pair studies, and were of Danish parentage for at least the prior three generations. The controls were collected at the University of Aarhus, and included 500 medical students, all of Danish parentage for at least three generations. All subjects gave written informed consent and the Danish Data Protection Agency and the ethics committees of Denmark approved the human subjects protocol.

1304

1305 Bipolar Disorder

#### 1306 Adolfsson, R | Not published | Umeå, Sweden | bip\_ume4\_eur

1307 Clinical characterization of the patients included the Mini-International Neuropsychiatric 1308 Interview (MINI), the Diagnostic Interview for Genetic Studies (DIGS), the Family Interview for 1309 Genetic Studies (FIGS) and the Schedules for Clinical Assessment in Neuropsychiatry (SCAN). 1310 The final diagnoses were made according to the DSM-IV-TR and determined by consensus of 2 1311 research psychiatrists. The unrelated Swedish control individuals, consisting of a large population-1312 based sample representative of the general population of the region, were randomly selected from 1313 the 'Betula study'.

#### 1314 Alda, M; Smoller, J | Not published | Nova Scotia, Canada; I2B2 controls | bip\_hal2\_eur

1315 The case samples were recruited from patients longitudinally followed at specialty mood disorders 1316 clinics in Halifax and Ottawa (Canada). Cases were interviewed in a blind fashion with the 1317 Schedule of Affective Disorders and Schizophrenia-Lifetime version (SADS-L) and consensus 1318 diagnoses were made according to DSM-IV and Research Diagnostic Criteria (RDC). Protocols 1319 and procedures were approved by the local Ethics Committees and written informed consent was 1320 obtained from all patients before participation in the study. Control subjects were drawn from the 1321 I2B2 (Informatics for Integrating Biology and the Bedside) project. The study consists of de-1322 identified healthy individuals recruited from a healthcare system in the Boston, MA, US area. The 1323 de-identification process meant that the Massachusetts General Hospital Institutional Review 1324 Board elected to waive the requirement of seeking informed consent as detailed by US Code of 1325 Federal Regulations, Title 45, Part 46, Section 116 (46.116).

#### Andreassen, OA | PMID:21926972 [PGC1], PMID:20451256 | Norway (TOP) | bip\_top7\_eur 1327 In the TOP study (Tematisk omrade psykoser), cases of European ancestry, born in Norway, were 1328 recruited from psychiatric hospitals in the Oslo region. Patients were diagnosed according to the 1329 SCID and further ascertainment details have been reported. Healthy control subjects were 1330 randomly selected from statistical records of persons from the same catchment area as the patient 1331 groups. The control subjects were screened by interview and with the Primary Care Evaluation of 1332 Mental Disorders (PRIME-MD). None of the control subjects had a history of moderate/severe 1333 head injury, neurological disorder, mental retardation or an age outside the age range of 18-60 1334 years. Healthy subjects were excluded if they or any of their close relatives had a lifetime history 1335 of a severe psychiatric disorder. All participants provided written informed consent and the human 1336 subjects protocol was approved by the Norwegian Scientific-Ethical Committee and the

1326

1337 Norwegian Data Protection Agency.

#### 1338 Andreassen, OA | Not published | Norway (TOP) | bip\_top8\_eur

1339 The TOP8 bipolar disorder cases and controls were ascertained in the same way as the 1340 bip top7 eur (TOP7) samples described above, and recruited from hospitals across Norway.

### 1341 Biernacka, JM; Frye, MA | 27769005 | Mayo Clinic, USA | bip may1 eur

1342 Bipolar cases were drawn from the Mayo Clinic Bipolar Biobank. Enrolment sites included Mayo 1343 Clinic, Rochester, Minnesota; Lindner Center of HOPE/University of Cincinnati College of 1344 Medicine, Cincinnati, Ohio; and the University of Minnesota, Minneapolis, Minnesota. Enrolment 1345 at each site was approved by the local Institutional Review Board approval, and all participants 1346 consented to use of their data for future genetic studies. Participants were identified through routine 1347 clinical appointments, from in-patients admitted in mood disorder units, and recruitment 1348 advertising. Participants were required to be between 18 and 80 years old and be able to speak 1349 English, provide informed consent, and have DSM-IV-TR diagnostic confirmation of type 1 or 2 1350 bipolar disorder or schizoaffective bipolar disorder as determined using the SCID. Controls were 1351 selected from the Mayo Clinic Biobank. Potential controls with ICD9 codes for bipolar disorder, 1352 schizophrenia or related diagnoses in their electronic medical record were excluded.

#### 1353 Blackwood, D | 18711365 [PGC1] | Edinburgh, UK | bip\_edi1\_eur

This sample comprised Caucasian individuals contacted through the inpatient and outpatient services of hospitals in South East Scotland. A BD-I diagnosis was based on an interview with the patient using the SADS-L supplemented by case note review and frequently by information from medical staff, relatives and caregivers. Final diagnoses, based on DSM-IV criteria were reached by consensus between two trained psychiatrists. Ethnically-matched controls from the same region were recruited through the South of Scotland Blood Transfusion Service. Controls were not directly screened to exclude those with a personal or family history of psychiatric illness. The study was approved by the Multi-Centre Research Ethics Committee for Scotland and patients gave written informed consent for the collection of DNA samples for use in genetic studies.

#### 1363 Breen, G; Vincent, JB | 24387768; 19416921; 21926972 [PGC1] |London, UK; Toronto,

#### 1364 **Canada [BACC] | bip\_bac1\_eur**

1365 The total case/control cohort (N=1922) includes 871 subjects from Toronto, Canada (N=431 cases 1366 (160 male; 271 female); N=440 controls (176 male; 264 female)), 1051 subjects from London, UK 1367 (N=538 cases (180 male; 358 female); N=513 controls (192 male; 321 female)). A summary of 1368 mean and median age at interview, age of onset (AOO), diagnostic subtypes (BD 1 versus BD 2), 1369 presence of psychotic symptoms, suicide attempt and family history of psychiatric disorders has 1370 been provided previously for both the Toronto and London cohorts. From the Toronto site (Centre 1371 for Addiction & Mental Health (CAMH)), BD individuals and unrelated healthy controls matched 1372 for age, gender and ethnicity were recruited. Inclusion criteria for patients: a) diagnosed with 1373 DSMIV/ICD 10 BD 1 or 2; b) 18 years old or over; c) Caucasian, of Northern and Western 1374 European origin, and three out of four grandparents also N.W. European Caucasian. Exclusion 1375 criteria include: a) Use of intravenous drugs; b) Evidence of intellectual disability; c) Related to 1376 an individual already in the study; d) Manias that only ever occurred in relation to or resulting 1377 from alcohol or substance abuse/dependence, or medical illness; e) Manias resulting from non-1378 psychotropic substance usage. The SCAN interview (Schedule for Clinical Assessments in 1379 Neuropsychiatry) was used for subject assessment. Using the SCAN interview along with case 1380 note review, each case was assigned DSM-IV and ICD 10 diagnoses by two independent 1381 diagnosticians, according to lifetime consensus best-estimate diagnosis. Lifetime occurrence of 1382 psychiatric symptoms was also recorded using the OPCRIT checklist, modified for use with mood

disorders. Similar methods and criteria were also used to collect a sample of 538 BD cases and
513 controls for the London cohort (King's College London; KCL). Both studies were approved
by respective institutional research ethics committees (the CAMH Research Ethics Board (REB)
in Toronto, and the College Research Ethics Committee (CREC) at KCL), and informed written
consent was obtained from all participants. GWAS results have previously been published for the
entire KCL/CAMH cohort.

#### 1389 Corvin, A | 18711365 [PGC1] | Ireland | bip\_dub1\_eur

1390 Samples were collected as part of a larger study of the genetics of psychotic disorders in the 1391 Republic of Ireland, under protocols approved by the relevant IRBs and with written informed 1392 consent that permitted repository use. Cases were recruited from Hospitals and Community 1393 psychiatric facilities in Ireland by a psychiatrist or psychiatric nurse trained to use the SCID. 1394 Diagnosis was based on the structured interview supplemented by case note review and collateral 1395 history where available. All diagnoses were reviewed by an independent reviewer. Controls were 1396 ascertained with informed consent from the Irish GeneBank and represented blood donors who 1397 met the same ethnicity criteria as cases. Controls were not specifically screened for psychiatric 1398 illness.

# 1399 Rietschel, M; Nöthen, MM, Cichon, S | 21926972 [PGC1] | BOMA-Germany I | 1400 bip\_bonn\_eur

1401 Cases for the BOMA-Bipolar Study were ascertained from consecutive admissions to the inpatient 1402 units of the Department of Psychiatry and Psychotherapy at the University of Bonn and at the 1403 Central Institute for Mental Health in Mannheim, University of Heidelberg, Germany. DSM-IV 1404 lifetime diagnoses of bipolar I disorder were assigned using a consensus best-estimate procedure, 1405 based on all available information, including a structured interview with the SCID and SADS-L, 1406 medical records, and the family history method. In addition, the OPCRIT checklist was used for 1407 the detailed polydiagnostic documentation of symptoms. Controls were ascertained from three 1408 population-based studies in Germany (PopGen, KORA, and Heinz-Nixdorf-Recall Study). The 1409 control subjects were not screened for mental illness. Study protocols were reviewed and approved 1410 in advance by Institutional Review Boards of the participating institutions. All subjects provided 1411 written informed consent.

## 1412 Rietschel, M; Nöthen, MM; Schulze, TG; Reif, A; Forstner, AJ | 24618891 | BOMA-Germany 1413 II | bip\_bmg2\_eur

Cases were recruited from consecutive admissions to psychiatric in-patient units at the University Hospital Würzburg. All cases received a lifetime diagnosis of BD according to the DSM-IV criteria using a consensus best-estimate procedure based on all available information, including semistructured diagnostic interviews using the Association for Methodology and Documentation in Psychiatry, medical records and the family history method. In addition, the OPCRIT system was used for the detailed polydiagnostic documentation of symptoms.

1420 Control subjects were ascertained from the population-based Heinz Nixdorf Recall (HNR) Study.
1421 The controls were not screened for a history of mental illness. Study protocols were reviewed and
1422 approved in advance by Institutional Review Boards of the participating institutions. All subjects
1423 provided written informed consent.

# 1424 Rietschel, M; Nöthen, MM; Schulze, TG; Bauer, M; Forstner, AJ; Müller-Myhsok, B | 1425 24618891 | BOMA-Germany III | bip\_bmg3\_eur

Cases were recruited at the Central Institute of Mental Health in Mannheim, University of
Heidelberg, and other collaborating psychiatric hospitals in Germany. All cases received a lifetime

1428 diagnosis of BD according to the DSM-IV criteria using a consensus best-estimate procedure based

1429 on all available information including structured diagnostic interviews using the AMDP, 1430 Composite International Diagnostic Screener (CID-S), SADS-L and/or SCID, medical records, 1431 and the family history method. In addition, the OPCRIT system was used for the detailed 1432 polydiagnostic documentation of symptoms. Controls were selected randomly from a Munich-1433 based community sample and recruited at the Max-Planck Institute of Psychiatry. They were 1434 screened for the presence of anxiety and mood disorders using the CID-S. Only individuals without 1435 mood and anxiety disorders were collected as controls. Study protocols were reviewed and 1436 approved in advance by Institutional Review Boards of the participating institutions. All subjects 1437 provided written informed consent.

#### 1438 Hauser, J; Lissowska, J; Forstner, AJ | 24618891 | BOMA-Poland | bip\_bmpo\_eur

1439 Cases were recruited at the Department of Psychiatry, Poznan University of Medical Sciences, 1440 Poznan, Poland. All cases received a lifetime diagnosis of BD according to the DSM-IV criteria 1441 on the basis of a consensus best-estimate procedure and structured diagnostic interviews using the 1442 SCID. Controls were drawn from a population-based case-control sample recruited by the Cancer-1443 Center and Institute of Oncology, Warsaw, Poland and a hospital-based case-control sample 1444 recruited by the Nofer Institute of Occupational Medicine, Lodz, Poland. The Polish controls were 1445 produced by the International Agency for Research on Cancer (IARC) and the Centre National de 1446 Génotypage (CNG) GWAS Initiative for a study of upper aerodigestive tract cancers. The controls 1447 were not screened for a history of mental illness. Study protocols were reviewed and approved in 1448 advance by Institutional Review Boards of the participating institutions. All subjects provided 1449 written informed consent.

#### 1450 Rietschel, M; Nöthen, MM; Rivas, F; Mayoral, F; Kogevinas, M; others | 24618891 | BOMA-

1451 Spain | bip\_bmsp\_eur

1452 Cases were recruited at the mental health departments of the following five centers in Andalusia, 1453 Spain: University Hospital Reina Sofia of Córdoba, Provincial Hospital of Jaen; Hospital of Jerez 1454 de la Frontera (Cádiz); Hospital of Puerto Real (Cádiz); Hospital Punta Europa of Algeciras 1455 (Cádiz); and Hospital Universitario San Cecilio (Granada). Diagnostic assessment was performed 1456 using the SADS-L; the OPCRIT; a review of medical records; and interviews with first and/or 1457 second degree family members using the Family Informant Schedule and Criteria (FISC). 1458 Consensus best estimate BD diagnoses were assigned by two or more independent senior 1459 psychiatrists and/or psychologists, and according to the RDC, and the DSM-IV. Controls were 1460 Spanish subjects drawn from a cohort of individuals recruited in the framework of the European 1461 Community Respiratory Health Survey (ECRHS, http://www.ecrhs.org/). The controls were not 1462 screened for a history of mental illness. Study protocols were reviewed and approved in advance 1463 by Institutional Review Boards of the participating institutions. All subjects provided written 1464 informed consent.

# Fullerton, J.M.; Mitchell, P.B.; Schofield, P.R.; Martin N.G.; Cichon, S. | 24618891 | BOMAAustralia | bip\_bmau\_eur

1467 Cases were recruited at the Mood Disorder Unit, Prince of Wales Hospital in Sydney. All cases 1468 received a lifetime diagnosis of BD according to the DSM-IV criteria on the basis of a consensus 1469 best-estimate procedure and structured diagnostic interviews using the DIGS, FIGS, and the SCID. 1470 Controls were parents of unselected adolescent twins from the Brisbane Longitudinal Twin Study. 1471 The controls were not screened for a history of mental illness. Study protocols were reviewed and 1472 approved in advance by Institutional Review Boards of the participating institutions. All subjects 1473 provided written informed consent.

#### 1474 Grigoroiu-Serbanescu, M; Nöthen, MM | 21353194 | BOMA-Romania | bip\_rom3\_eur

1475 Cases were recruited from consecutive admissions to the Obregia Clinical Psychiatric Hospital, 1476 Bucharest. Patients were administered the DIGS and FIGS interviews. Information was also 1477 obtained from medical records and close relatives. The diagnosis of BP-I was assigned according 1478 to DSM-IV criteria using the best estimate procedure. All patients had at least two hospitalized 1479 illness episodes. Population-based controls were evaluated using the DIGS to exclude a lifetime 1480 history of major affective disorders, schizophrenia, schizoaffective disorders, and other psychoses, 1481 obsessive-compulsive disorder, eating disorders, and alcohol or drug addiction.

#### 1482 Craddock, N, Jones, I, Jones, L | 17554300 | WTCCC | bip\_wtcc\_eur\_sr-qc

1483 Cases were all over the age of 17 yr, living in the UK and of European descent. Recruitment was 1484 undertaken throughout the UK and included individuals who had been in contact with mental 1485 health services and had a lifetime history of high mood. After providing written informed consent, 1486 participants were interviewed by a trained psychologist or psychiatrist using a semi-structured 1487 lifetime diagnostic psychiatric interview (Schedules for Clinical Assessment in Neuropsychiatry) 1488 and available psychiatric medical records were reviewed. Using all available data, best-estimate 1489 life-time diagnoses were made according to the RDC. In the current study we included cases with 1490 a lifetime diagnosis of RDC bipolar 1 disorder, bipolar 2 disorder or schizo-affective disorder, 1491 bipolar type. Controls were recruited from two sources: the 1958 Birth Cohort study and the UK 1492 Blood Service (blood donors) and were not screened for history of mental illness. All cases and 1493 controls were recruited under protocols approved by the appropriate IRBs. All subjects gave 1494 written informed consent.

#### 1495 Kelsoe, J | 21926972 [PGC1] | USA (GAIN) | bip\_gain\_eur

1496 *Genetic Association Information Network (GAIN)/ The Bipolar Genome Study (BiGS)* The BD 1497 sample was collected under the auspices of the NIMH Genetics Initiative for BD 1498 (http://zork.wustl.edu/nimh/), genotyped as part of GAIN and analyzed as part of a larger GWAS 1499 conducted by the BiGS consortium. Approximately half of the GAIN sample was collected as 1500 multiplex families or sib pair families (waves 1-4), the remainder were collected as individual 1501 cases (wave 5). Subjects were ascertained at 11 sites: Indiana University, John Hopkins University, 1502 the NIMH Intramural Research Program, Washington University at St. Louis, University of 1503 Pennsylvania, University of Chicago, Rush Medical School, University of Iowa, University of 1504 California, San Diego, University of California, San Francisco, and University of Michigan. All 1505 investigations were carried out after the review of protocols by the IRB at each participating 1506 institution. At all sites, potential cases were identified from screening admissions to local treatment 1507 facilities and through publicity programs or advocacy groups. Potential cases were evaluated using 1508 the DIGS, FIGS, and information from relatives and medical records. All information was 1509 reviewed through a best estimate diagnostic procedure by two independent and non-interviewing 1510 clinicians and a consensus best-estimate diagnosis was reached. In the event of a disagreement, a 1511 third review was done to break the tie. Controls were from the NIMH Genetic Repository sample 1512 obtained by Dr. P. Gejman through a contract to Knowledge Networks, Inc. Only individuals with 1513 complete or near-complete psychiatric questionnaire data who did not fulfill diagnostic criteria for 1514 major depression and denied a history of psychosis or BD were included as controls for BiGS 1515 analyses. Controls were matched for gender and ethnicity to the cases.

1516 Kelsoe, J; Sklar, P; Smoller, J | [PGC1 Replication] | USA (FAT2; FaST, BiGS, TGEN) |
1517 bip\_fat2\_eur

Cases were collected from individuals at the 11 U.S. sites described for the GAIN sample. Eligible participants were age 18 or older meeting DSM-IV criteria for BD-I or BD-II by consensus diagnosis based on interviews with the Affective Disorders Evaluation (ADE) and MINI. All participants provided written informed consent and the study protocol was approved by IRBs at
each site. Collection of phenotypic data and DNA samples were supported by NIMH grants
MH063445 (JW Smoller); MH067288 (PI: P Sklar), and MH63420 (PI: V Nimgaonkar). The
control samples were NIMH controls that were using the methods described in that section. The
case and control samples were independent of those included in the GAIN sample.

#### 1526 Kirov, G | 25055870 | Bulgarian trios | bip\_butr\_eur

All cases were recruited in Bulgaria from psychiatric inpatient and outpatient services. Each proband had a history of hospitalisation and was interviewed with an abbreviated version of the SCAN. Consensus best-estimate diagnoses were made according to DSM-IV criteria by two researchers. All participants gave written informed consent and the study was approved by local ethics committees at the participating centers.

#### 1532 Kirov, G | 25055870 | UK trios | bip\_uktr\_eur

The BD subjects were recruited from lithium clinics and interviewed in person by a senior psychiatrist, using abbreviated version of the SCAN. Consensus best-estimate diagnoses were made based on the interview and hospital notes. Ethics committee approval for the study was obtained from the relevant research ethics committees and all individuals provided written informed consent for participation.

#### 1538 Landén, M; Sullivan, PF; Sklar, P | [ICCBD] | Sweden (ICCBD) | bip\_swa2\_eur

The BD subjects were identified using the Swedish National Quality Register for Bipolar Disorders (BipoläR) and the Swedish National Patient Register (using a validated algorithm requiring at least two hospitalizations with a BD diagnosis). A confirmatory telephone interview with a diagnostic review was conducted. Additional subjects were recruited from the St. Göran Bipolar Project (Affective Center at Northern Stockholm Psychiatry Clinic, Sweden), enrolling new and ongoing 1544 patients diagnosed with BD using structured clinical interviews. Diagnoses were made according

1545 to the DSM-IV criteria (BipoläR and St. Göran Bipolar Project) and ICD-10 (National Patient

1546 Register). The control subjects used were the same as for the SCZ analyses described above. All

ascertainment procedures were approved by the Regional Ethical Committees in Sweden.

#### 1548 Landén, M; Sullivan, PF; Sklar, P | [ICCBD] | Sweden (ICCBD) | bip\_swei\_eur

1549 The cases and controls in the bip\_swei\_eur sample were recruited using the same ascertainment 1550 methods described for the bip\_swa2\_eur sample.

#### 1551 Leboyer, M | [PGC1 replication] | France | bip\_fran\_eur

Cases with BD1 or BD2 and control samples were recruited as part of a large study of genetics of BD in France (Paris-Creteil, Bordeaux, Nancy) with a protocol approved by relevant IRBs and with written informed consent. Cases were of French descent for more than 3 generations were assessed by a trained psychiatrist or psychologist using structured interviews supplemented by medical case notes, mood scales and self-rating questionnaire assessing dimensions.

#### 1557 Li, Q | 24166486; 27769005 | USA (Janssen), SAGE controls | bip\_jst5\_eur

1558 The study included unrelated patients with bipolar 1 disorder from 6 clinical trials (IDs: 1559 NCT00257075, NCT00253162, NCT00076115, NCT00299715, NCT00309699, and 1560 NCT00309686). Participant recruitment was conducted by Janssen Research & Development, 1561 LLC (formerly known as Johnson & Johnson Pharmaceutical Research & Development, LLC) to 1562 assess the efficacy and safety of risperidone. Bipolar cases were diagnosed according to DSM-IV-1563 TR criteria. The diagnosis of bipolar disorder was confirmed by the Schedule for Affective 1564 Disorders and Schizophrenia for School-Age Children-Present and Lifetime Version (K-SADS-PL) in NCT00076115, by the SCID in NCT00257075 and NCT00253162, or by the MINI in 1565 1566 NCT00299715 and NCT00309699, and NCT00309686, respectively. Additional detailed descriptions of these clinical trials can be found at ClinicalTrials.gov. Only patients of European
ancestry with matching controls were included in the current analysis. Controls subjects were
drawn from the Study of Addiction: Genetics and Environment (SAGE, dbGaP Study Accession:
phs000092.v1.p1). Control subjects did not have alcohol dependence or drug dependence
diagnoses; however, mood disorders were not an exclusion criterion.

## McQuillin, A; Gurling, H | 18317468 [PGC1] | UCL (University College London), London, UK | bip\_uclo\_eur

1574 The UCL sample comprised Caucasian individuals who were ascertained and received clinical 1575 diagnoses of bipolar 1 disorder according to UK National Health Service (NHS) psychiatrists at 1576 interview using the categories of the International Classification of Disease version 10. In addition 1577 bipolar subjects were included only if both parents were of English, Irish, Welsh or Scottish 1578 descent and if three out of four grandparents were of the same descent. All volunteers read an 1579 information sheet approved by the Metropolitan Medical Research Ethics Committee who also 1580 approved the project for all NHS hospitals. Written informed consent was obtained from each 1581 volunteer. The UCL control subjects were recruited from London branches of the National Blood 1582 Service, from local NHS family doctor clinics and from university student volunteers. All control 1583 subjects were interviewed with the SADS-L to exclude all psychiatric disorders.

## 1584 Craddock, N; Jones, I; Jones, L | [ICCBD] | Cardiff and Worcester, UK (ICCBD-BDRN) | 1585 bip\_icuk\_eur

Cases were all over the age of 17 yr, living in the UK and of European descent. Cases were recruited via systematic and not systematic methods as part of the Bipolar Disorder Research Network project (<u>www.bdrn.org</u>), provided written informed consent and were interviewed using a semi-structured diagnostic interview, the Schedules for Clinical Assessment in Neuropsychiatry. 1590 Based on the information gathered from the interview and case notes review, best-estimate lifetime 1591 diagnosis was made according to DSM-IV. Inter-rater reliability was formally assessed using 20 1592 randomly selected cases (mean  $\kappa$  Statistic = 0.85). In the current study we included cases with a 1593 lifetime diagnosis of DSM-IV bipolar disorder or schizo-affective disorder, bipolar type. The 1594 BDRN study has UK National Health Service (NHS) Research Ethics Committee approval and 1595 local Research and Development approval in all participating NHS Trusts/Health Boards.Controls 1596 were part of the Wellcome Trust Case Control Consortium common control set, which comprised 1597 healthy blood donors recruited from the UK Blood Service and samples from the 1958 British 1598 Birth Cohort. Controls were not screened for a history of mental illness. All cases and controls 1599 were recruited under protocols approved by the appropriate IRBs. All subjects gave written 1600 informed consent.

#### 1601 **Ophoff, RA | Not Published | Netherlands | bip\_ucla\_eur**

The case sample consisted of inpatients and outpatients recruited through psychiatric hospitals and institutions throughout the Netherlands. Cases with DSM-IV bipolar disorder, determined after interview with the SCID, were included in the analysis. Controls were collected in parallel at different sites in the Netherlands and were volunteers with no psychiatric history after screening with the (MINI). Ethical approval was provided by UCLA and local ethics committees and all participants gave written informed consent.

#### 1608 Paciga, S | [PGC1] | USA (Pfizer) | bip\_pf1e\_eur

This sample comprised Caucasian individuals recruited into one of three Geodon (ziprasidone) clinical trials (NCT00141271, NCT00282464, NCT00483548). Subjects were diagnosed by a clinician with a primary diagnosis of Bipolar 1 Disorder, most recent episode depressed, with or without rapid cycling, without psychotic features, as defined in the DSM-IV-TR (296.5x) and 1613 confirmed by the MINI (version 5.0.0). Subjects also were assessed as having a HAM-D-17 total 1614 score of >20 at the screening visit. The trials were conducted in accordance with the protocols, 1615 International Conference on Harmonization of Good Clinical Practice Guidelines, and applicable 1616 local regulatory requirements and laws. Patients gave written informed consent for the collection 1617 of blood samples for DNA for use in genetic studies.

#### 1618 Pato, C | [ICCBD] | Los Angeles, USA (ICCBD-GPC)| bip\_usc2\_eur

Genomic Psychiatry Consortium (GPC) cases and controls were collected via the University of Southern California healthcare system, as previously described. Using a combination of focused, direct interviews and data extraction from medical records, diagnoses were established using the OPCRIT and were based on DSM-IV-TR criteria. Age and gender-matched controls were ascertained from the University of Southern California health system and assessed using a validated screening instrument and medical records.

# Scott, L; Myer, RM; Boehnke, M | 19416921 [PGC1] | Michigan, USA (Pritzker and NIMH) | bip\_mich\_eur

1627 The Pritzker Neuropsychiatric Disorders Research Consortium (NIMH/Pritzker) case and controls 1628 samples were from the NIMH Genetics Initiative Genetics Initiative Repository. Cases were 1629 diagnosed according to DMS-III or DSM-IV criteria using diagnostic interviews and/or medical 1630 record review. Cases with low confidence diagnoses were excluded. From each wave 1-5 available 1631 non-Ashkenazi European-origin family, two BD1 siblings were included when possible and the 1632 proband was preferentially included if available (n=946 individuals in 473 sibling pairs); otherwise 1633 a single BD1 case was included (n=184). The bipolar sibling pairs were retained within the 1634 NIMH/Pritzker sample when individuals in more than one study were uniquely assigned to a study 1635 set. Controls had non-Ashkenazi European-origin, were aged 20-70 years and reported no diagnosis with or treatment for BD or schizophrenia, and that they had not heard voices that others
could not hear. Individuals with suspected major depression were excluded based on answers to
questions related to depressive mood. NIMH controls were further selected as the best match(es)

1639 to NIMH cases based on self-reported ancestry.

#### 1640 Sklar, P; Smoller, J | 18317468 [PGC1] | USA (STEP1) | bip\_stp1\_eur

1641 The Systematic Treatment Enhancement Program for Bipolar Disorder (STEP-BD) was a seven-1642 site, national U.S., longitudinal cohort study designed to examine the effectiveness of treatments 1643 and their impact on the course of BD that enrolled 4,361 participants who met DSM-IV criteria 1644 for BD1, BD2, bipolar not otherwise specified (NOS), schizoaffective manic or bipolar type, or 1645 cyclothymic disorder based on diagnostic interviews. From the parent study, 2,089 individuals 1646 who were over 18 years of age with BD1 and BD2 diagnoses consented to the collection of blood 1647 samples for DNA. BD samples with a consensus diagnosis of BD1 were selected for inclusion in 1648 STEP1. Two groups of controls samples from the NIMH repository were used. One comprised 1649 DNA samples derived from US Caucasian anonymous cord blood donors. The second were 1650 controls who completed the online self-administered psychiatric screen and were ascertained as 1651 described above, by Knowledge Networks Inc. For the second sample of controls only those 1652 without history of schizophrenia, psychosis, BD or major depression with functional impairment 1653 were used.

#### 1654 Sklar, P; Smoller, J | 18711365 [PGC1] | USA (STEP2) | bip\_stp2\_eur

The STEP2 sample included BD-1 and BD-2 samples from the STEP-BD study described above along with BD-2 subjects from UCL study also described above. The controls samples for this study were from the NIMH repository as described above for the STEP1 study.

1658

#### 1659 European ancestry, trio design

1660 Schizophrenia

#### 1661 Kirov, G: Owen M | 22083728 Bulgaria | ms.scz\_butr\_eur

1662 Families from Bulgaria were recruited if a proband had schizophrenia or schizoaffective disorder, 1663 both parents were available, and all members of the trio agreed to participate in the study. 1664 Recruitment took place between 1999 and 2004 in several psychiatric hospitals in Bulgaria. Ethical 1665 Committee approval was obtained from each of these hospitals. All probands and all parents 1666 received an Information Sheet and signed Informed Consent Forms. All participants had attended 1667 mainstream schools, which at the time in Bulgaria, excluded people with mental retardation. 1668 Probands were either in- or out-patients at the time of the study but each had a history of 1669 hospitalization. A team of psychiatrists was trained in using the rating scales and methods of the 1670 study. We used the SCAN instrument to perform an interview for psychotic and mood symptoms. 1671 This instrument has been translated into Bulgarian and validated by one of its authors (A. 1672 Jablensky). Consensus diagnoses were made according to DSM-IV criteria on the basis of an 1673 interview and inspection of hospital notes by two clinicians. If consensus was not attained, the 1674 patient was re-interviewed by a research interview trained clinician and was excluded if consensus 1675 could still not be reached. In addition, approximately 23% of the sample was selected at random 1676 and re-interviewed by a research interview trained clinician. Hospital notes were also collected for 1677 affected relatives in order to confirm diagnoses.

1678 Levinson, D | 22885689 | Six countries | ms.scz\_lemu\_eur

1679 Schizophrenia cases were included from the family sample of European-ancestry pedigrees 1680 described by Levinson et al. Participants and their families in this trio study, probands were 1681 ascertained and recruited from different clinical settings (e.g. inpatients, outpatients and 1682 community facilities) in six countries (Australia, France, Germany, Ireland, UK, and the US). 1683 (Unrelated individuals were included as part of a case-control design, see Levinson, D, 1684 scz lacw eur above.) Diagnoses were established using semi-structured interviews, psychiatric 1685 records and informant reports. Case probands were diagnosed with schizophrenia or 1686 schizoaffective disorder according to DSM-III-R criteria. The trio-based analysis included families 1687 where there was at least one affected proband and two available parents. Each affected sibling in 1688 such families was included, with the parents, as an independent trio. All protocols were approved 1689 by loci IRBs, and all cases provided written informed consent.

#### 1690 Kirov, G: Owen, M | Not Published | Bulgaria | ms.scz\_uktr\_eur

All cases and parents were recruited from UK and had a history of hospitalization for treatment of schizophrenia. Diagnosis was confirmed following a SCAN interview and review of case notes followed by consensus diagnosis according to DSM-IV criteria. The samples were genotyped at the Broad Institute. All participants gave written informed consent and the study was approved by local ethics committees at the participating centers. The samples were genotyped at the Broad Institute.

1697

#### 1698 Genotype Quality Control

To ensure independence of the data sets, individuals were excluded until no individual showed a relatedness (pihat) value greater than 0.2 to any other individual in the collection, while preferentially keeping the case over the control for case-control related pairs. In total 1,795 BD cases, 1,165 SCZ cases and 27,274 controls were removed (most of which were previously known), leaving 20,129 BD cases 33,426 SCZ cases and 54,065 controls for the final metaanalysis. 1705 For analyses directly comparing BD and SCZ, we matched cases from both phenotypes on 1706 genotyping platform and ancestry, resulting in 15,270 BD cases versus 23,585 SCZ cases. Hence, 1707 we were able to match 76% of BD cases and 71% of SCZ cases for this case vs case analysis. 1708 Among our entire dataset, 44% of the sample was female, 51% was male and 5% were unreported

by the collection site. This work focused explicitly on the autosomes and sought maximal power 1710 across the analyses, sex was not used except for during quality control and sex-specific analyses 1711 were not performed in this effort. Individual ages were not provided. For a subset of cases, we had

1712 information for age of onset which were used in subphenotype specific analyses only.

1713

1709

#### 1714 **Sub-phenotype Description**

1715 BD sub-phenotypes were collected by each study site using a combination of diagnostic 1716 instruments, case records and participant interviews. Ascertainment details for each study site are 1717 described in the supplementary data of the PGC Bipolar Working Group paper(Stahl et al., 2017). 1718 The selection of phenotypes for collection by this group was determined by literature searches in 1719 order to determine phenotypes with prior evidence for heritability. It was further refined dependent 1720 on the availability of phenotype data across a range of study sites and the consistency by which 1721 the phenotypes were defined. Schizophrenia subphenotypes represent quantitative traits extracted 1722 using factor analysis from a set of standard psychiatric assessments and represent four symptom 1723 dimensions (manic, depressive, positive and negative). These subphenotypes were used 1724 previously(Ruderfer et al., 2014) but in this work we have increased the sample size with additional 1725 cohorts being added.

1726

#### 1727 **METHOD DETAILS**

1728

#### 1729 QUANTIFICATION AND STATISTICAL ANALYSIS

1730

#### 1731 Quality Control, Imputation, Association Analysis and Polygenic Risk Score Testing

1732 Quality control and imputation were performed on each of the study cohort datasets (n=81), 1733 according to standards established by the Psychiatric Genomics Consortium (PGC). The quality 1734 control parameters for retaining SNPs and subjects were: SNP missingness < 0.05 (before sample removal); subject missingness (p < 0.02); autosomal heterozygosity deviation ( $|F_{het}| < 0.2$ ); SNP 1735 1736 missingness < 0.02 (after sample removal); difference in SNP missingness between cases and controls < 0.02; and SNP Hardy-Weinberg equilibrium ( $p > 10^{-6}$  in controls or  $p > 10^{-10}$  in cases). 1737 1738 Genotype imputation was performed using the pre-phasing/imputation stepwise approach 1739 implemented in IMPUTE2(Howie et al., 2011) / SHAPEIT(Delaneau et al., 2013) (chunk size of 1740 3 Mb and default parameters). The imputation reference set consisted of 2,186 phased haplotypes 1741 from the full 1000 Genomes Project dataset (August 2012, 30,069,288 variants, release 1742 "v3.macGT1"), all variants align to human genome build 19 (hg19). After imputation, we used the 1743 best guess genotypes (genotype probability > 0.8), for further robust relatedness testing and 1744 population structure analysis. Here we required very high imputation quality (INFO > 0.8) and low missingness (<1%) for further quality control. After linkage disequilibrium (LD) pruning ( $r^2$  < 1745 1746 0.02) and frequency filtering (MAF > 0.05), there were 14,473 autosomal SNPs in the data set. 1747 Principal component estimation was done with the same collection of autosomal SNPs. We tested 1748 the first 20 principal components for phenotype association (using logistic regression with study 1749 indicator variables included as covariates) and evaluated their impact on the genome-wide test 1750 statistics using  $\lambda$ . Thirteen principal components namely 1,2,3,4,5,6,7,8,10,12,15,18,20 were

included in all association analyses ( $\lambda$ =1.45). Analytical steps were repeated for SCZ vs BD analysis.

1753 We performed four main association analyses (Figure 1), i.e. (i) GWAS of BD and SCZ as a single 1754 combined case phenotype, as well as disorder-specific GWAS using independent control sets in 1755 (ii) BD cases vs BD controls and (iii) SCZ cases vs SCZ controls, and (iv) association analysis of 1756 SCZ cases vs BD cases. For all GWS loci from the GWAS of BD and SCZ vs controls we identified 1757 any GWS loci within 1Mb from the extent of the locus in the previously published PGC SCZ vs 1758 controls(Schizophrenia Working Group of the Psychiatric Genomics Consortium, 2014) and the 1759 most recent PGC GWAS of BD vs controls(Stahl et al., 2017) and performed conditional analysis. 1760 Specifically, we transformed the genotype probabilities of the disease variant into dosages and 1761 used it as an additional covariate for the association analysis for the BD+SCZ vs controls index 1762 SNP. This was done within each cohort and an OR based inverse SE weighted meta-analysis was 1763 performed for the final result. All datasets were included except for those with trios.

1764

#### 1765 Summary-data-based Mendelian Randomization (SMR)

1766 SMR(Zhu et al., 2016) is a method that integrates summary level GWAS data with gene expression 1767 quantitative trait loci (eQTL) identified in independent data sets. This integration aims to identify 1768 variants that have pleotropic effects on expression of a given gene and the phenotype. While 1769 significant findings may indeed reflect a causal path from variant to phenotype through expression, 1770 it is impossible to discern statistically between pleiotropy and causality. However, the method can 1771 remove linkage as driving the result, and uses the data available to prioritise amongst genes in 1772 genomic regions that show association with disease. We used SMR as a statistical fine-mapping 1773 tool applied to the SCZ vs BD GWAS results to identify loci with strong evidence of causality via 1774 gene expression. SMR analysis is limited to significant (FDR < 0.05) cis SNP-expression 1775 quantitative trait loci (eQTLs) with MAF > 0.01. eQTLs passing these thresholds were combined 1776 with GWAS results in the SMR test, with significance (p<sub>SMR</sub>) reported at a Bonferroni-corrected 1777 threshold for each eQTL data set. The eQTL architecture may differ between genes. For example, 1778 through LD, many SNPs can generate significant associations with the same gene, but in some 1779 instances multiple SNPs may be independently associated with the expression of a gene. After 1780 identification of significant SNP-expression-trait association through the SMR test, a follow-up 1781 heterogeneity test aims to prioritize variants by excluding regions for which there is conservative 1782 evidence for multiple causal loci ( $p_{HET} < 0.05$ ). SMR analyses were conducted using eQTL data 1783 from whole peripheral blood (Westra et al., 2013), dorsolateral prefrontal cortex generated by the CommonMind Consortium<sup>8</sup>, and 11 brain sub-regions from the GTEx consortium(Consortium, 1784 1785 2015).

1786

#### 1787 **Regional joint GWAS**

1788 Summary statistic Z-scores were calculated for each marker in each of the four main GWAS 1789 results, using the logistic regression coefficient and its standard error. Rare SNPs (MAF < 0.01), 1790 and SNPs with a low INFO score (< 0.3) in either dataset were removed. The causal variant 1791 relationships between SCZ and BD were investigated using the Bayesian method software pw-1792 gwas (v0.2.1), with quasi-independent regions determined by estimate LD blocks in an analysis of 1793 European individuals (n=1,703)(Berisa and Pickrell, 2015; Pickrell et al., 2016). Briefly, pw-gwas 1794 takes a Bayesian approach to determine the probability of five independent models of association. 1795 (1) There is no causal variant in BD or SCZ; (2) a causal variant in BD, but not SCZ (3); a causal 1796 variant in SCZ, but not BD; (4) a shared causal variant influencing both BD and SCZ; (5) two

causal variants where one influences BD, and one influences SCZ (Figure 2). The posterior
probability of each model is calculated using model priors, estimated empirically within pw-gwas.
Regions were considered to support a particular model when the posterior probability of the model
was greater than 0.5.

1801

#### 1802 **Regional SNP-heritability estimation**

1803 We calculated local SNP-heritability independently for SCZ and BD using the Heritability 1804 Estimator from Summary Statistics (HESS) software(Shi et al., 2016) for each of the independent 1805 regions defined above. The sum of these regional estimates is the total SNP-heritability of the trait. 1806 To calculate local SNP-heritability HESS requires reference LD matrices representative of the 1807 population from which the GWAS samples were drawn. We utilized the 1000 genomes European 1808 individuals as the reference panel (The 1000 Genomes Project Consortium, 2015). Unlike pw-1809 gwas(Pickrell et al., 2016), HESS does not assume that only one causal variant can be present in 1810 each region.

1811

#### 1812 DATA AND SOFTWARE AVAILABILITY

1813 Summary statistics from GWAS are publically available at https://www.med.unc.edu/pgc/results-1814 and-downloads/downloads.

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