Copy number variants encompassing Mendelian disease genes in a large multigenerational family segregating bipolar disorder

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Abstract
Background: Bipolar affective disorder (BP) is a common, highly heritable psychiatric disorder characterized by periods of depression and mania. Using dense SNP genotype data, we characterized CNVs in 388 members of an Old Order Amish Pedigree with bipolar disorder. We identified CNV regions arising from common ancestral mutations by utilizing the pedigree information. By combining this analysis with whole genome sequence data in the same individuals, we also explored the role of compound heterozygosity.

Results: Here we describe 541 inherited CNV regions, of which 268 are rare in a control population of European origin but present in a large number of Amish individuals. In addition, we highlight a set of CNVs found at higher frequencies in BP individuals, and within genes known to play a role in human development and disease. As in prior reports, we find no evidence for an increased burden of CNVs in BP individuals, but we report a trend towards a higher burden of CNVs in known Mendelian disease loci in bipolar individuals (BPI and BPII, p = 0.06).

Conclusions: We conclude that CNVs may be contributing factors in the phenotypic presentation of mood disorders and co-morbid medical conditions in this family. These results reinforce the hypothesis of a complex genetic architecture underlying BP disorder, and suggest that the role of CNVs should continue to be investigated in BP data sets.

Keywords: CNV, Bipolar disorder, Family based studies, Mendelian disease genes, Genetics loci

Background
Bipolar affective disorder (BP) is a serious mental disorder characterized by periodic changes in mood, energy and activity levels alternating between episodes of depression and mania [1]. The lifetime prevalence of BP type I (BPI) and type II (BPII) is 2.1% in the United States [2] and the age of onset is early, at 18–19.5 years old [3], making BP responsible for the loss of more disability-adjusted life-years than all forms of cancer [4] and consequently it is a major public health concern [5]. As with many complex disorders, the underlying etiology of BP is unknown, but is hypothesized to be the result of multiple gene-gene and gene-environment interactions [6]. Epidemiological studies using twin data show that BP has heritability estimates ranging from 62-89% [7,8], although the mode of inheritance is complex. Common genetic factors have been shown to contribute substantially to susceptibility for bipolar disorder, with a strong polygenic contribution [9]. Several potential BP candidate genes have been described [10], but findings are inconsistent and the role of specific genes in BP is currently undetermined.

Copy number polymorphisms (CNVs) are a common class of genetic variation in the human genome [11-14], and can be readily detected using intensity data from genome-wide SNP arrays. Like single-nucleotide polymorphisms (SNPs), CNVs can affect gene expression, either by encompassing genes or regulatory elements. Large, cytogenetically detectable chromosomal rearrangements, such as aneuploidy, have been historically linked to human disease [15]. Studies of several genomic disorders, associated with inherited or sporadic genomic anomalies which are...
smaller (<5 MB) and therefore can not be detected using conventional cytogenetic methods, revealed that deletions and duplications encompassing several genes may lead to complex and highly pleiotropic clinical syndromes [16].

Several systematic surveys of copy number variation using Comparative Genomic Hybridization (CGH) and high-density SNP arrays revealed a large number of benign deletions and duplications across the genome, but also revealed the role of a large number of rare potentially pathogenic CNVs in neurodevelopmental and psychiatric diseases, particularly in Autism Spectrum Disorder and Schizophrenia [17]. In Autism Spectrum Disorder, CNVs were found in a number of chromosomal regions [18], and the burden of rare and de novo CNVs was enriched in affected individuals compared to controls and their unaffected siblings [19]. Similarly, several large, rare CNVs have been associated with schizophrenia [20,21]. Among these CNVs several have been observed at elevated rates in multiple neurodevelopmental and psychiatric disorders [22,23].

Both linkage and candidate gene analyses, as well as genome-wide association studies, indicate a shared genetic architecture and an overlap of susceptibility between BP and schizophrenia [24]. However, compared to studies conducted on ASD and schizophrenia, there are far fewer examples of CNVs associated with BP [25]. An analysis of 1001 cases and 1034 controls reported an increased burden of singleton CNVs in early onset bipolar cases [26]. Also, in an independent study of 788 trios, frequencies of de novo CNVs were significantly higher in bipolar disorder as compared to controls, but not as high as in schizophrenia [27]. However, a study using Welcome Trust Case Control Consortium (WTCCC) data found no evidence for an elevated burden of CNVs in bipolar individuals (n = 1697) compared to controls (n = 2806), although the burden was found to be elevated in schizophrenia [28]. The same authors recently published the most comprehensive analysis of CNVs in the WTCCC revealing a significantly lower rate of rare very large CNVs (>1 Mb) in patients with bipolar disorder (n = 1,650) compared to reference individuals without psychiatric disorder (n = 10,259) [29]. Although the authors state that this result needs to be verified in larger datasets, they propose that a lower CNV burden may underlie differences in the presentation of clinical phenotype between bipolar disorder and schizophrenia. In addition, recent research suggests that de novo CNVs may play a smaller role in BP compared to schizophrenia [30], but the role of inherited CNVs remains uncertain.

The Old Order Amish are a founder population originating in middle Europe. Since 1964, when Victor McKusick and colleagues described the benefits from medical genetics studies in the Amish [31], a large number of Mendelian disorders have been described in this population [32]. More recently, next generation sequencing studies of neurodevelopmental and psychiatric disorders in the Amish provide a unique opportunity to address the role of rarer forms of genetic variation [33,34]. However, these recent studies focus on the role of single nucleotide variants (SNVs). Apart from a handful of gene deletions associated with Mendelian disease [32], and 50 CNV regions identified in a subset of individuals from the Old Order Amish pedigree with bipolar disorder [35], global analysis of copy number variation has not been systematically carried out in this genetic isolate.

The aim of the present study was to investigate CNVs in the extended Old Order Amish pedigree with bipolar disorder, and compare these CNVs with CNVs detected in a large collection of unrelated control subjects to identify deletions and duplications private to this family. Also, we compared burden and frequency of CNVs in family members with affective disorders (BPI, BPII and MDD-R) with their unaffected relatives to identify CNVs potentially contributing to the locus and allele heterogeneity of bipolar disorder. Our systematic analysis revealed 67 rare and moderately rare CNVs encompassing Mendelian disease genes that may contribute to the complex and pleiotropic manifestation of mental illness in this founder population.

Results

Overall strategy

To characterize structural variants in 388 members of a large multigenerational Old Order Amish pedigree with bipolar disorder, we used dense SNP genotype data generated using the Illumina Omni 2.5 M platform [33]. We also performed CNV analysis on 2,156 Age-related Eye Disease Study (AREDS) control subjects (1,897 with European ethnicity) genotyped using the same SNP platform. A flowchart (Figure 1) outlines the quality control and analysis pipeline employed to address: a) differences in the allele frequency of CNVs in this genetic isolate compared to a large sample of subjects of European origin; b) the role of CNVs in susceptibility to bipolar illness in this large pedigree; c) an estimate of the total per genome (or person) burden of CNVs, including CNVs that encompass known disease loci.

A catalog of inherited CNVs in an Amish pedigree segregating bipolar disorder

As part of our genetic study of bipolar disorder in the Old Order Amish, we analyzed genome-wide SNP genotype data using the Penn CNV algorithm [36] to identify CNVs. We examined the breakpoints of all CNVs (n = 18,986) and clustered groups of CNVs that have arisen from common ancestral mutations (see Methods) into CNV regions (n = 561). Using the pedigree relationships, we classified all regions as either ‘inherited’ (shown to pass from parent to child), or of ‘unknown origin’ (not seen in either parent). To avoid possible technical artifacts in the analysis, we focused on the high quality, inherited CNVs observed.
in multiple (more than three) Amish family members. These variants are less likely to represent false positives in computational prediction or rearrangements that arose during culturing of lymphoblastoid cell lines [37]. Furthermore, in a large pedigree with an excess of bipolar disorder, we expect that the causal genetic variants will be inherited, rather than de novo. Of 541 inherited CNV regions identified by the analysis of 328 family members (a subset that contains both parent and child information), 33 overlap with 50 regions previously identified in a small scale study of the core family, i.e. 50 family members [35]. Eight CNV regions (four which are exonic, one which is intronic, and three which are intergenic) were detected on Chromosome X (Additional file 1: Table S2).

Among detected inherited CNV regions, the largest category consisted of common CNVs (present in more than 5% of controls) found throughout the large multigenerational Amish pedigree. In addition we detected 104 moderately rare (present in less than 5% of controls) and 139 rare (present in less than 1% of controls) CNV regions in subjects, as well as 129 ‘exclusive’ regions that were not present in any controls. Of these exclusive regions, 36 are deletions and 93 are duplications, and 99 regions include genes (Figure 2, created using Circos [38]). These ‘exclusive’ variants form a key part of the genomic architecture of this pedigree, and could play a role in phenotypic presentation. To illustrate the frequency of a CNV in the pedigree, CNV counts are presented for nuclear families only in which the CNV is present, and only for those individuals with bipolar or well phenotypes; individuals with unknown or other phenotypes are excluded. They include a 26 kb duplication on 13q24, present in 109 Amish individuals (affected 23/86, 26.4%; unaffected 66/232, 28.4%), which encompasses the entire SRY (sex determining region Y)-box 1 (SOX1) gene; a 24 kb deletion on 5q33.1, found in 48 individuals (affected 8/38, 21.1%; unaffected 29/79, 36.7%), in an intergenic region upstream of both coiled-coil domain containing 69 (CCDC69) and GM2 ganglioside activator (GM2A); and a 10 kb deletion on 12q21.31, found in 33 individuals (affected 9/25, 36.0%; unaffected 19/51, 37.3%), located downstream of solute carrier family 6, member 15 (SLC6A15).

We compared the total number, average size, and burden per individual of CNVs in a) all family members (n = 375), b) subjects with bipolar disorder (n = 77), and unaffected Amish and control individuals (Amish n = 234, controls n = 1897) (Table 1). Analysis was performed on copy number losses (deletions) and gains (duplications) separately. Overall number of detected deletions and
duplications in the Amish pedigree were comparable to those detected in the control subjects.

**CNVs and disease association**

No individual CNV segregated fully with bipolar disorder. Analysis of CNV data in the linkage regions previously reported [33] identified a single duplication event in the 7q21 region. The 95 kb duplication localizes upstream of the maximum LOD score marker D7S518. We confirmed inheritance of the CNV on the 4-4-1-4 haplotype (D7S2431-D7S554-D7S18-D7S2509). The duplication spans the first exon of the collagen, type XXVI, alpha 1 (COL26A1) gene. COL26A1 has yet to be functionally characterized, with a possible role in aspirin-intolerant asthma [39].

Burden analysis of CNV regions in genes in the Amish shows a trend towards an increased number of these
Table 1 Summary of CNV calls for the Old Order Amish (n = 375) and controls (n = 1897)

|                          | Total Deletions | Total Duplications | Average Size (bp) | Burden (per individual) |
|--------------------------|-----------------|--------------------|-------------------|--------------------------|
|                          | Amish CNVs      | Amish Inherited CNVs | Controls | Amish CNVs | Amish Inherited CNVs | Controls | Amish CNVs | Amish Inherited CNVs | Controls |
| Total Number             |                 |                    |                   |                          |                  |
| All Samples              | 18986           | 6345               | 77205             | 10942                    | 4154             | 51138     | 8044         | 2191          | 26067     |
| Affected                 | 3844            | 1380               | -                 | 2216                     | 905              | -         | 1628         | 475           | -         |
| Unaffected               | 12191           | 3962               | -                 | 6022                     | 2581             | -         | 5269         | 1381          | -         |
| Average Size (bp)        |                 |                    |                   |                          |                  |           |              |               |           |
| All Samples              | 36123           | 42064              | 29697             | 31124                    | 32701            | 23768     | 42924        | 59815         | 41328     |
| Affected                 | 41123           | 44572              | -                 | 39802                    | 33575            | -         | 42921        | 65523         | -         |
| Unaffected               | 34363           | 39932              | -                 | 29072                    | 32093            | -         | 41315        | 54581         | -         |
| Burden (per individual)  |                 |                    |                   |                          |                  |           |              |               |           |
| All Samples              | 50.6            | 23.2               | 40.7              | 29.2                     | 15.2             | 27.0      | 21.6         | 8.2           | 13.7      |
| Affected                 | 49.9            | 23.4               | -                 | 28.8                     | 15.3             | -         | 21.1         | 8.1           | -         |
| Unaffected               | 52.1            | 23.7               | -                 | 29.6                     | 15.5             | -         | 22.5         | 8.3           | -         |

Total number, average size, and burden of CNVs was calculated for all Amish CNVs, inherited Amish CNVs, and control CNVs. CNVs were analyzed together, and as deletion and duplication events separately.
CNVs in bipolar individuals (narrow phenotype: BPI and BPII), although this does not reach significance (narrow burden: 17.3, unaffected burden: 15.6, p = 0.11) (Table 2). We identified three rare deletions in KCNJ6, LINC13C, OTOL1 and 7 rare duplications in CNTNAP2/MIR548F3, CORO7/VASN, DTNB, EMID2, KCNFI, PDPR and SGTA1/THOP1 that are present in children with bipolar disorder (and their parents). In addition, we find other rare CNVs in genes that are present frequently in individuals with bipolar disorder (Table 3). Association analysis for all CNV regions was performed using two different methods: a) FBAT [40] and b) EMMAX [41], although no CNV was found to be significantly associated with BP following correction for multiple testing. We found no overall enrichment of large inherited CNVs in affected individuals, although 7 large, rare, CNVs in genes occurred more frequently in subjects with bipolar disorder than unaffected family members and control subjects. One of the largest rare genic CNVs is the previously reported 150 kb deletion in the 15q11 region, which encompasses the entire Prader-Willi region non-protein coding RNA 2 (PWRN2) gene [35]. The deletion is present in 15 families, is found on two haplotypes (D15S817-D15S1021-D15S128: 3-6-3 and 3-5-3) and is widely spread throughout the pedigree; 20/32 (62.5%) of those with bipolar disorder in these carrier families have the CNV, compared to 28/52 (53.8%) of well individuals.

Next, we focused on the analysis of CNVs encompassing known disease genes. It has been suggested that heterozygosity for several mutations in Mendelian disease genes may lead to complex disease risk, such as behavioral anomalies in neurodevelopmental and psychiatric disorders [63]. To ask if CNVs in disease genes may contribute to the allelic architecture in the Amish family segregating bipolar disorder, we mapped known disease genes. It has been suggested that heterozygosity for several mutations in Mendelian disease genes may lead to complex disease risk, such as behavioral anomalies in neurodevelopmental and psychiatric disorders [63]. To ask if CNVs in disease genes may contribute to the allelic architecture in the Amish family segregating bipolar disorder, we mapped known disease genes. We identified three rare deletions in CORO7/VASN, DTNB, EMID2, KCNFI, PDPR and SGTA1/THOP1 that are present in children with bipolar disorder (and their parents). In addition, we find other rare CNVs in genes that are present frequently in individuals with bipolar disorder (Table 3). Association analysis for all CNV regions was performed using two different methods: a) FBAT [40] and b) EMMAX [41], although no CNV was found to be significantly associated with BP following correction for multiple testing. We found no overall enrichment of large inherited CNVs in affected individuals, although 7 large, rare, CNVs in genes occurred more frequently in subjects with bipolar disorder than unaffected family members and control subjects. One of the largest rare genic CNVs is the previously reported 150 kb deletion in the 15q11 region, which encompasses the entire Prader-Willi region non-protein coding RNA 2 (PWRN2) gene [35]. The deletion is present in 15 families, is found on two haplotypes (D15S817-D15S1021-D15S128: 3-6-3 and 3-5-3) and is widely spread throughout the pedigree; 20/32 (62.5%) of those with bipolar disorder in these carrier families have the CNV, compared to 28/52 (53.8%) of well individuals.

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Table 2 Burden analysis of CNV Regions reveals a higher number of CNVs in genes in individuals with narrow bipolar phenotype (BPI and BPII)

|                   | Unaffected | Broad | Narrow |
|-------------------|------------|-------|--------|
| No. All CNVs      | 28.1       | 28.7  | 29.5   |
| P vs unaffected   | -          | 0.39  | 0.26   |
| No. All CNVs in genes | 15.6       | 16.8  | 17.3   |
| P vs unaffected   | -          | 0.16  | 0.11   |
| No. Rare CNVs in genes | 9.7        | 11.0  | 11.3   |
| P vs unaffected   | -          | 0.12  | 0.09   |
| No. CNVs in disease genes | 4.8        | 5.5   | 5.7    |
| P vs unaffected   | -          | 0.11  | 0.06   |
| No. Rare CNVs in disease genes | 4.0        | 4.8   | 5.0    |
| P vs unaffected   | -          | 0.07  | 0.06   |

A trend towards an increased number of CNVs in disease genes in individuals with narrow bipolar phenotype is also reported.
| Cytoband | Start | Stop | No. snp | Length (bp) | CN | Contained genes | Frequency | No. subjects | No. affected | No. families | Predicted effect | FBAT | EMMAX | Previous disease associations for gene |
|----------|-------|------|---------|-------------|----|----------------|-----------|--------------|-------------|-------------|----------------|------|-------|--------------------------------------|
| 1p36.21  | 13171723 | 13218942 | 31 | 47220 | 0&1 | LOC440563 | Rare | 38 | 13 | 15 | Gene del | 1 | 0.1887 |  |
| 1q22     | 155152005 | 155162067 | 22 | 13287 | 3 | MUC1, TRIM46 | Amish specific | 28 | 13 | 32 | Partial gene dup | 0.0711 | 0.1535 | Kidney disease [42] |
| 1q24.1   | 165644865 | 165649715 | 9 | 4851 | 1 | ALODH9A1 | Very rare | 17 | 7 | 12 | Exonic del | 1 | 0.1434 |  |
| 2q31.1   | 176929113 | 177000696 | 86 | 71584 | 3 | EVX2, HOXD10, HOXD11, HOXD12, HOXD13, HOXD8, HOXD9 | Amish specific | 45 | 15 | 32 | Gene dup | 0.4122 | 0.9834 | Limb and genital abnormalities [43,44] |
| 2q37.3   | 241500674 | 241516094 | 20 | 24920 | 3 | DUSP28, RNPEPL1 | Very rare | 74 | 13 | 43 | Partial gene dup | 0.4226 | 0.3229 |  |
| 2q37.3   | 241482099 | 241516094 | 20 | 24920 | 3 | ANKMY1, DUSP28, RNPEPL1 | Very rare | 31 | 7 | 23 | Partial gene dup for ANKMY1 and RNPEPL1, gene dup for DUSP28 | 0.0007 | 0.3229 |  |
| 3p21.2   | 51989546 | 51995419 | 11 | 10143 | 3 | GPR62, PCBP4 | Amish specific | 72 | 17 | 47 | Partial gene dup | 0.0412 | 0.0238 |  |
| 3p25.3   | 11411823 | 11414339 | 5 | 2517 | 0&1 | ATG7 | Rare | 46 | 13 | 24 | Intrinsic del | 0.7557 | 0.7020 | Frontotemporal dementia, Parkinsons disease [45] |
| 3q29     | 193136358 | 193140348 | 9 | 3991 | 0&1 | ATP13A4 | Rare | 41 | 9 | 18 | Intrinsic del | 0.7389 | 0.2519 | Autism [46] |
| 4q22.1   | 91907363 | 91913329 | 16 | 10710 | 1 | FAM190A | Amish specific | 34 | 6 | 21 | Intrinsic del | 0.0099 | 0.0794 |  |
| 5q35.3   | 179221537 | 179238794 | 25 | 32170 | 3 | LTC4S, MGAT4B, MIR1229, SQSTM1, MAML1 | Amish specific | 79 | 17 | 36 | Partial gene dup for LTC4S and SQSTM1, gene dup for MGAT4B and MIR1229 | 0.6641 | 0.0973 | Venous thromboembolism and ischaemic stroke [47], Paget disease of bone [48] |
| 5q35.3   | 179211629 | 179231681 | 25 | 32170 | 3 | LTC4S, MGAT4B, MIR1229 | Amish specific | 61 | 14 | 36 | Gene dup | 0.2513 | 0.0973 | Venous thromboembolism and ischaemic stroke [47] |
| 6p21.32  | 32610719 | 32614917 | 10 | 4199 | 1 | HLA-DQA1 | Rare | 25 | 7 | 13 | Exonic del | 1 | 0.0574 |  |
| 6p25.3   | 1612234 | 1620037 | 15 | 9536 | 3 | FOXC1 | Very rare | 47 | 14 | 23 | Exonic del | 1 | 0.5815 | Axenfeld-Rieger anomaly [49] |
| 6q26     | 163041460 | 163139315 | 64 | 99376 | 1 | PARK2 | Very rare | 21 | 5 | 11 | Exonic del | 0.5637 | 0.5534 | Parkinsons disease [50], Autism [51] |
| 7q22.1   | 100968058 | 101063059 | 159 | 183210 | 3 | EMID2 | Rare | 100 | 25 | 38 | Exonic dup | 0.0330 | 0.7031 |  |
| 7q36.1   | 149461487 | 149516968 | 67 | 55482 | 3 | SSPO, ZNF467 | Amish specific | 154 | 39 | 68 | Partial gene dup | 0.8907 | 0.1100 |  |
| 8p21.3   | 21943602 | 22024523 | 80 | 62524 | 3 | FAM160B2, HR, NUDT18 | Amish specific | 112 | 20 | 59 | Partial gene dup for FAM160B2 and HR, gene dup for NUDT18 | 0.0284 | 0.9422 | Alopecia universalis [52], Congenital Atrichia [53] |
| 8p22     | 15947559 | 16023673 | 118 | 76115 | 1 | MSR1 | Rare | 19 | 3 | 7 | Partial gene del | 0.0833 | 0.6317 | Prostate cancer [54] |
| Chromosome | Start | End | Length | Number | Disease | Gene | Disease Specific | Common Gene | FBAT EMMAX p-values | Reference |
|------------|-------|-----|--------|--------|----------|------|----------------|-------------|-----------------|-----------|
| 8p22       | 15419777 | 15432653 | 15432653 | 21108 | 1 | TUSC3 | Very rare | Intronic del | 0.0833 | 0.6317 | Intellectual disability [55] |
| 9q34.11    | 130497180 | 130518716 | 24 | 21108 | TOR2A | Very rare | SH2D3C | Partial gene dup for TOR2A | 0.0116 | 0.4779 |
| 9q34.11    | 45359483 | 45222200 | 125 | 151274 | Rare | TMEM72-AS1 | Partial gene dup | 0.4386 | 0.5237 |
| 10q21.3    | 68239474 | 68422442 | 109 | 152110 | Very rare | CTNNA3 | Partial gene del | 0.6547 | 0.5957 | Arrhythmogenic right ventricular cardiomyopathy [56] |
| 11p11.2    | 45916436 | 45931646 | 24 | 29093 | 3 | C11orf94, MAPK8IP1, PEX16 | Amish specific | Partial gene dup for MAPK8IP1 and PEX16, gene dup for C11orf94 | 0.0197 | 0.0402 | Diabetes type 2 [57], Zellweger syndrome [58] |
| 11p15.4    | 8959020 | 8964938 | 11 | 5919 | Very rare | ASC1 | Gene del | 0.4795 | 0.3681 |
| 13q34      | 112712459 | 112726336 | 26 | 26199 | Very rare | SOX1 | Gene dup | 0.0254 | 0.0749 | Neuronal development [59] |
| 13q34      | 114518789 | 114530659 | 31 | 17552 | Very rare | GAT6 | Partial gene dup | 0.6946 | 0.6193 |
| 14q23.2    | 63957653 | 63962909 | 5 | 6398 | Very rare | PPP2R5E | Intronic del | 0.2568 | 0.1877 |
| 15q11.2    | 24345146 | 24496990 | 76 | 152110 | Rare | PPWRN2 | Gene del | 0.1967 | 0.4189 | Prader-Willi region [60] |
| 15q26.1    | 90615898 | 90636762 | 28 | 26809 | Amish specific | IDH2, ZNF710 | Partial gene dup | 0.7389 | 0.3569 | D-2-hydroxyglutaric aciduria, type II [61] |
| 16p12.1    | 27337036 | 27350687 | 15 | 20228 | Very rare | IL4R | Exonic del | 0.7812 | 0.1062 |
| 17p13.3    | 811982 | 1183612 | 665 | 456481 | Very rare | ABR, BHLH9, MIR3183, 10XN, TIMM22, TUSC5 | Partial gene dup for 10XN and TUSC5, gene dup for ABR, BHLH9, MIR3183 and TIMM22 | 0.6547 | 0.3319 |
| 18q23      | 77150335 | 77162816 | 40 | 24952 | Rare | NFATC1 | Amish specific | Exonic dup | 0.8273 | 0.1415 | Tricuspid atresia [62] |
| 18q23      | 76725624 | 76767375 | 35 | 41752 | Very rare | SALL3 | Gene dup | 0.1336 | 0.0864 |
| 21q22.3    | 44822871 | 44868905 | 35 | 46025 | Rare | SIK1 | Exonic dup | 0.3938 | 0.1313 |

CNVs shown are rare in controls (present in fewer than 5% of controls. Rare: <5%, Very rare: <1%, Amish Specific: not found in controls), and common in the Amish (present in more than 5% of individuals). Contained genes show all genes in CNV, disease genes are highlighted in bold. FBAT and EMMAX p-values for association analysis for Bipolar disorder are included, p<0.05 are in bold.
permitted the investigation of a combined effect of CNVs with likely deleterious SNPs on the same and in trans haplotype. Table 4 lists 26 disease genes with both a CNV and at least one non-synonymous SNP present in the same gene in the same individual. Among CNVs in known disease genes are two rare CNVs at the PARK2 locus: a 63 kb deletion and a 99 kb deletion spanning the second exon. Although these CNVs do not segregate with a bipolar disease status, it is notable that 300 kb distal to the CNV breakpoint, in the neighboring PARK2 co-regulated gene (PACRG), we previously detected a cluster of SNPs with a family-based association signal of p-value 2.16x10-6 for the top SNP (rs9365506) (Additional file 1: Figure S3) [33]. In addition, there are six individuals who have both an exonic missense variant (rs1801582) and a CNV in PARK2. The variant is located upstream of the CNV and in each individual is present on a different haplotype from the CNV. Haplotype analysis of the region shows multiple haplotypes containing the SNPs from the family-based association analysis, two of which contain the CNVs, further supporting a proposed clustering of several potential risk alleles (SNPs and CNVs) in a defined chromosomal region [33].

Also among these variants was a rare 2 kb intronic deletion in autophagy related 7 (ATG7), which is present in 46 individuals, out of which 13 have a bipolar phenotype (13/36, 36.1%; unaffected 25/124, 20.2%). Furthermore, individuals with this CNV also have a possibly damaging exonic missense SNP (rs36117895) present on the same haplotype as the CNV. We also identified a number of individuals (n = 19, affected 3/6, 50%; unaffected 14/39, 35.9%) with a rare CNV in the intron of tumor suppressor candidate 3 (TUSC3). These individuals carry a SNP (rs1035972) within the same gene on the same haplotype as the CNV. Although many of these individuals are ‘unaffected’ (14 unaffected, 3 affected) within our pedigree, mutations such as these may contribute to the overall burden of disease within the Amish population.

**Table 4 Compound heterozygosity - Disease genes with CNVs and variants in the same individual**

| Gene       | Number of affected (Total = 69) | Number of unaffected (Total = 203) | CNV frequency |
|------------|---------------------------------|-----------------------------------|---------------|
| ATG7       | 13                              | 26                                | Rare          |
| SNTG1      | 4                               | 2                                 | Common        |
| PTPRD      | 4                               | 3                                 | Rare          |
| IL4R       | 9                               | 20                                | Rare          |
| HLA-DQA1   | 7                               | 15                                | Rare          |
| KCNQ6      | 1                               | 0                                 | Amish Specific|
| CCDC50     | 6                               | 15                                | Common        |
| Dicer1     | 1                               | 1                                 | Amish Specific|
| GALNTL4    | 1                               | 1                                 | Rare          |
| ATP13A4    | 9                               | 25                                | Rare          |
| ERBB4      | 1                               | 3                                 | Common        |
| Msr1       | 1                               | 3                                 | Rare          |
| Park2      | 1                               | 3                                 | Rare          |
| RHD        | 2                               | 6                                 | Common        |
| CDH13      | 0                               | 1                                 | Rare          |
| PrkG1      | 0                               | 1                                 | Rare          |
| WwDX       | 3                               | 10                                | Common        |
| Smarca2    | 2                               | 8                                 | Common        |
| Cyp2D6     | 0                               | 3                                 | Rare          |
| Tusc3      | 3                               | 14                                | Rare          |
| UgT1A7     | 0                               | 7                                 | Rare          |
| UgT1A8     | 0                               | 7                                 | Rare          |
| UgT1A10    | 0                               | 8                                 | Rare          |
| UgT1A13    | 0                               | 8                                 | Rare          |
| Ctnina3    | 0                               | 9                                 | Rare          |
| Cacna1c    | 0                               | 14                                | Common        |

Counts in each column represent the number of individuals with both a CNV and another variant in the same gene.

**Discussion**

We recently reported a combined analysis of dense genotypes and whole genome sequence for a large Old Order Amish pedigree with bipolar disorder. This study focused on the analysis of missense mutations within linkage peaks and detected a high degree of genetic heterogeneity of mental illness in this family [33]. Here we report results of the analysis of CNVs in the same extended pedigree. The mean burden per individual and size of CNVs were similar between our Amish sample and the European controls. While previous studies of the role of CNVs as risk alleles for bipolar disorder have been limited, an increased burden of CNVs in bipolar disorder has been reported by some [26,27], although these findings are not consistent [28,29]. We find a trend towards an increased burden of CNVs in genes in individuals with BP, specifically for CNVs that are rare or moderately frequent in the general population. While this finding does not reach significance, it adds to a body of evidence that suggests that CNVs may have some as-yet undefined role in BP and should be investigated accordingly.

We identified 13 CNVs in genes previously associated with psychiatric and developmental disorders, that are present at a higher rate in the Amish extended pedigree than in the control sample. The comparison of the frequency of these structural anomalies in the extended pedigree and in a large control dataset identified disease associated CNVs that are enriched in this population and may serve as a starting point for a future ‘Genotype-first
“approach” [66] to defining subtypes of bipolar and other complex diseases in this founder population. For example, rare CNVs were found in SOX1, a transcriptional activator thought to play a role in neuronal development [59]; and near GM2A, which is highly expressed in the brain and can harbor mutations which result in a variant of Tay-Sachs disease [67]; and SLC6A15, an amino acid transporter expressed highly in the brain and associated with major depressive disorder [68]. The largest rare, genic deletion in our pedigree, encompassing PWRN2 and the surrounding region, was found more frequently in bipolar individuals within carrier families. PWRN2 lies within a 1.5 Mb section on the long arm of chromosome 15 found to be deleted in Prader-Willi syndrome, a neurogenetic disorder with cognitive, behavioral and endocrine phenotypes [69]. A duplication in the HOXD region on chromosome 2 was also more present frequently in bipolar individuals. The HOXD genes play important roles in morphogenesis, and deletions in this cluster have been associated with limb and genital abnormalities [70].

Our study design allowed us to interrogate a combination of CNVs and other inherited mutations found within the same gene in a single individual on the same or opposite chromosome. Using this method, we identify 26 known disease genes that contain both a CNV and exonic mis-sense SNP in one or more individuals. Of these, particular genes of interest include ATG7, which has been associated with frontotemporal dementia [71]; TUSC3, a gene associated with intellectual disability [50]; and PARK2, a gene associated with Parkinson’s disease [55,72]. Although we do not provide evidence that these CNVs and SNPs alone are disease causing in this pedigree, they may contribute together with other variants within the same chromosomal region to the disease risk [33]. In addition, molecular studies of these reported CNVs would be needed to determine if they have any effect on the gene. Larger studies in a non-Amish population are also required to determine if CNVs at these loci could be of relevance to bipolar or psychiatric disorder in a general population.

Although clinical information for the large extended Old Order Amish pedigree is limited to mood disorders, our genetic data permits the analysis of CNVs in genes associated with Mendelian diseases. In our initial report on the analysis of CNVs in the core Amish pedigree (in 50 family members), we provided proof of principle for a family-based investigation of a combination of structural variants in the same subject as that could confer risk for a disease [35]. Our reported trend for an increased burden of disease CNVs in bipolar family members (when compared to their unaffected relatives) needs to be further investigated with a larger sample size, both in the founder and general population. In many Mendelian diseases, psychiatric and behavioral symptoms are prevalent [73] and a wide range of medical co-morbidities are common in psychiatric disorders [63]. The variants underlying Mendelian disease are generally highly penetrant and less influenced by the environment, while haploinsufficiency or heterozygosity at several Mendelian disease loci may lead to complex behavioral anomalies in psychiatric disorders. Moreover, such a burden of risk alleles could explain the high degree of heritability but rather complex genetic architecture observed in these disorders. In other words, we propose that some of the hidden heritability may reside in gene-by-gene interactions and that the analysis of interactions at bona fide disease genes may be well powered by focusing on the impact of genetic variation within these critical classes of genes.

We have identified over 100 CNVs with a significant difference in allele frequency between the Amish family and the control sample. These structural variants add to an extended list of non-synonymous, likely deleterious variants that are rare in the 1000 Genomes project dataset (<2%), but present in 1-30% of BP subjects and their family members [33]. Owing to the anonymized fashion in which our study was conducted, it will not be possible, at this time, to evaluate possible phenotypic consequences of private Amish structural variants, those that are present in >10% of family members in this pedigree, but rare or absent in the general population. However, several ongoing genetics research initiatives involving the Plain populations [33,34] combined with the clinical genetics profiling applied in several clinics that serve these communities, are generating valuable insights that could potentially allow prevention of disability and disease. As reported by the Clinic For Special Children and by colleagues involved in genetic studies in Hutterites, providing education and offering clinical carrier status for devastating Mendelian diseases would likely be welcomed by members of the founder communities [74,75].

Our study has multiple limitations. First of all, the analysis is focused on a large extended family and power is limited for the statistical assessment. Also, it is difficult to determine if CNVs found to be enriched in the pedigree are unique to this founder population or to a cluster of these families, originating primarily from the Lancaster area (in Pennsylvania). However, the use of a genetic isolate and a large family structure provided us with a higher level of genetic and phenotypic homogeneity, and permitted the tracing of CNV events within nuclear families and across generations. Also, although the availability of biomaterials through a public cell repository represents a major advantage of this collection, DNA isolated from lymphoblastoid cell lines rather than blood represents a limitation of our study. Therefore, to avoid possible cell line artifacts, we excluded all singleton CNVs from our analysis and we were not able to address the role of de novo CNVs. Other studies have
reported a role for de novo CNVs in BP [27], and we were unable to address this area of research. We note that we utilized CNVs from an eye disease study (AREDS) as a comparison for our dataset. Although we were not directly comparing the levels of CNVs between datasets, a disease free control dataset would have been more desirable. However, as their primary use was to identify CNVs that are present in the Amish more frequently than a European population, we believe the AREDS dataset was adequate for this purpose. Finally, the CNVs reported here were not experimentally validated, but due to our ability to show inheritance of the CNV from parent to child through the pedigree, we consider them to be validated CNVs [76].

Conclusions
In summary, we identify a number of CNVs in an Old Order Amish pedigree segregating BP. Many of the CNVs found were rare in the general population and present in a large number of Amish individuals. Some of the CNVs were found in a higher frequency in individuals with a BP phenotype, and within genes known to play a role in human development and disease. We conclude that these CNVs may be contributing factors in the phenotypic presentation and heterogeneity of mental illness in this family.

Methods
Sample
The genetic-epidemiologic study of bipolar disorder among the Old Order Amish in Pennsylvania (The Amish Study of Major Affective Disorder) has been well documented [77,78]. Diagnostic methods included structured interviews (SADS-L) that were conducted with the patients and close others. In addition, medical records were obtained following signed, informed consent, these were abstracted and collated for five members of the Psychiatric Board who were blind to patient names, pedigree, address, admission/discharge diagnosis and treatment. The Psychiatric Board members used strict Research Diagnostic Criteria (RDC) and the Diagnostic and Statistical Manual of Mental Disorders, 4th Edition (DSM-IV) for uniform clinical criteria, and reviewed all material every few years as a reliability check on diagnoses. The majority of affected individuals in the current pedigree are diagnosed as either BPI, BPII, or Major Depressive Disorder (MDD, recurrent) with a few Schizoaffective Disorder, BP Subtype, although there is a wider spectrum of Major Affective Disorders in the extended Amish pedigrees. In this study we place individuals into a number of phenotype categories for analysis: Narrow (BPI and BPII only), Broad (BPI, BPII and MDDR), and well (unaffected only).

Collection of blood samples followed diagnostic consensus. Lymphoblastoid cell lines were established by the Coriell Institute of Medical Research (CIMR). Signed informed consents were obtained to access medical records for the Amish Study clinicians exclusively to do diagnostic evaluations and clinical studies. Two forms were used: a) one with yearly Institutional Review Board (IRB, University of Miami) approval adhering to special guidelines because the Amish are defined as a “vulnerable” population; and b) a second using state approved, medical record consent forms for specific mental health clinics and psychiatric hospitals throughout central Pennsylvania. Collection of blood/tissue samples followed diagnostic consensus, using two informed consent forms: a) one with annual Univ. Miami IRB approval defining (with language appropriate for Old Order Amish) how their cells would be preserved for medical research on Major Affective Disorders; and, b) the Informed Consent Form required by the Institute for Medical Research (CIMR), later Coriell - National Institute for General Medical Sciences (NIGMS) Human Genetic Cell Repository (HGCR). In addition, analysis of whole-genome sequence data from consented individuals in this pedigree was also approved by the IRB of the Weill Cornell Medical College and the Perelman School of Medicine at the University of Pennsylvania.

Control subjects
Control subjects were selected from the Age-Related Eye Disease Study (AREDS) sponsored by the National Institutes of Health (National Eye Institute). This prospective study of about 3600 participants follows the clinical course of age-related macular degeneration (AMD) and age-related cataract. Participants in this study were required to be ‘free of any illness or condition that would make long-term follow-up or compliance with study medications unlikely or difficult’ and as such are considered ‘well’ for the purposes of our study of mental illness. In addition, age of participants recruited to this study was between 55 to 80 years old, beyond the age at which presentation of a bipolar phenotype is to be expected. The individuals studied here were not affected with macular degeneration or cataract at the AREDS baseline examination.

Collection of blood samples for genetic research was performed following recruitment. Lymphoblastoid cell lines were established by the Coriell Institute of Medical Research (CIMR). Genotyping was performed on 2159 AREDS samples using Illumina Omni 2.5 M SNP arrays at the Center for Inherited Disease Research (CIDR). We performed rigorous quality control of the raw genotype calls by applying a series of filters on both markers and samples using PLINK [79]. The initial dataset contained 2,443,179 SNPs and 2159 samples. The following filters were applied in sequence; the numbers of markers or samples excluded is given in parentheses: a) exclude SNPs with
missing rate > 0.5 (34,066), b) exclude samples with missing rate > 0.02 (0), c) exclude SNPs with missing rate > 0.02 (41,053), d) exclude SNPs with MAF < 0.02 (863,429), and e) exclude SNPs deviating from Hardy-Weinberg equilibrium at p < 1e-6 (34,405). After quality controls we retained 1,470,226 SNPs and 2159 samples. We subsequently performed Multidimensional Scaling on a set of ~500 k overlapping SNPs for all available AREDS genotypes and 1000 Genomes data. This analysis permitted the selection of 1897 subjects with European ancestry (Additional file 1: Figure S1) for further CNV calling and analysis.

Genotyping
Genotyping was performed on 394 samples from the extended Amish pedigree using Illumina Omni 2.5 M SNP arrays at the Center for Applied Genomics (Children’s Hospital of Pennsylvania, Philadelphia, PA). As with the AREDS data, we performed rigorous quality control of the raw genotype calls by applying a series of filters on both markers and samples using PLINK [79], with the addition of excluding markers based on informative missingness and individuals/markers based on the mendel error rate. The initial dataset contained 2,379,855 SNPs and 394 samples. The following filters were applied in sequence; the numbers of markers or samples excluded is given in parentheses: a) exclude SNPs with missing rate > 0.5 (19,435), b) exclude samples with missing rate > 0.02 (6), c) exclude SNPs with missing rate > 0.02 (31,678), d) exclude SNPs with MAF < 0.02 (1,018,805), e) exclude SNPs with informative missingness p < 1e-6 (0), f) exclude SNPs deviating from Hardy-Weinberg equilibrium at p < 1e-6 (0), g) exclude individuals with > 5% Mendelian errors (0) and h) exclude SNPs with >1% Mendelian errors (1334). After quality controls we retained 1,309,937 SNPs and 388 samples.

Association analysis for all CNV regions was performed using two different methods: a) FBAT [40] (Version 2.0.4), a version of the transmission distortion test adapted for larger families and b) EMMAX [41] (Version from February 2012), a statistical test for association analysis using mixed models that accounts for the population structure within the sample.

Identification of copy number variants (CNVs)
CNVs were called by PennCNV, a previously described CNV detection algorithm [36], using the GC model wave adjustment [80]. CNVs were removed if they had a value > 0.30 standard deviation of LRR (LRRSD), a waveness factor (WF) value > 0.05, or < 5 SNPs. Regions that are known to be highly unreliable for CNV calls, such as immunoglobulin regions and the centromeres/telomeres of chromosomes were excluded from the analysis (see Additional file 1: Table S1). Samples that had a total CNV number greater than 3 SD from the mean, or samples that showed evidence of aneuploidy, were also excluded. After quality control we retained a set of 18,986 CNVs in 375 individuals from the Amish sample, and 77,205 in 1,897 individuals from the AREDS sample.

Inherited CNV regions
From all available Amish samples with genotype data, we selected 328 individuals that belong to a nuclear family (parent plus children, 54 parents and 274 children) to ascertain regions containing inherited CNVs. This method consists of two stages. First, we establish the CNV region boundaries from the CNV that has the greatest overlap with other CNVs in the same genomic region. All CNVs that overlap 50% with this CNV are considered part of that CNV region. Second, we trace the inheritance of a CNV region using the pedigree information. For a CNV region to be inherited, it must be present in both the child and at least one parent.

Human disease catalog
The Human Genome Mutation Database (HGMD) catalogs known disease associated variants (http://www.hgmd.org/). Most of the clinical phenotypes in the database are monogenic diseases. In its most recent release (June 2013) it contains 141,000 different variants in ~5,700 genes (“HGMD disease genes”) [81]. We cataloged all CNV regions (detected by the analysis of dense genotypes for the 328 Amish family members) that partially or fully overlap 3457 HGMD disease genes (‘DM’ tag in HGMD).

Whole genome sequencing
Whole genome sequencing (WGS) for 80 Old Order Amish family members (including 30 parent child trios) was performed by Complete Genomics Inc. (CGI; Mountain View, CA) using a sequence-by-ligation method [82]. Paired-end reads of length 70 bp (35 bp at each end) were mapped to the National Center for Biotechnology Information (NCBI) human reference genome (build 37.2) using a Bayesian mapping pipeline [83]. Variant calls were performed by CGI using version 2.0.3.1 of their pipeline. False discovery rate estimates for SNP calls of the CGI platform are 0.2–0.6% [82]. Gene annotations were based on the NCBI build 37.2 seq_gene file contained in a NCBI annotation build. The variant calls within the WGS were processed using the cgatools software (version 1.5.0, build 31) made available by CGI. The listvar tool was used to generate a master list of the 11.1 M variants present in the 80 Amish samples. The testvar tool was used to determine presence and absence of each variant within the 80 Amish WGS. Only variants with high variant call scores (“VQHIGH” tag in the data files) were included. For further QC measures see [33].
As described in Georgi et al. [33], we performed phasing and imputation of variants identified by WGS into the Omni 2.5 M SNP genotypes using the Genotype Imputation Given Inheritance (GIGI) software version 1.02. GIGI performs imputation of dense genotypes in large pedigrees based on a sparse panel of framework markers using a Markov Chain Monte Carlo approach. Overall performance of our imputation is comparable to the published report [84]. For a threshold on the genotype imputation posterior probability of 0.85, we observed overall concordance of ~0.96 with a call rate of ~0.50. As expected, imputation performance increases for sub-pedigrees with a higher number of samples with WGS, i.e., when considering only nuclear families with WGS samples the performance improves to concordance ~0.99 and call rate ~0.87 [33].

**Additional file**

Additional file 1: Figures and Tables illustrating quality control measures, and supplementary results including additional CNV regions found in disease genes and on Chromosome X.

**Competing interests**

The authors declare that they have no competing interests.

**Authors’ contributions**

RK carried out the CNV analysis and drafted the manuscript. BG participated in the Amish project and carried out the whole genome sequence analysis. JEBW contributed the ARED’s data and gave advice for the analysis. DS contributed the ARED’s data and gave advice for the analysis. SMP and MB participated in the design of the study. MB also conceived the study and helped to draft the manuscript. All authors read and approved the final manuscript.

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