RESEARCH ARTICLE

Alu element in the RNA binding motif protein, X-linked 2 (RBMX2) gene found to be linked to bipolar disorder

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Abstract

Objective
We have used long-read single molecule, real-time (SMRT) sequencing to fully characterize a ~12Mb genomic region on chromosome Xq24-q27, significantly linked to bipolar disorder (BD) in an extended family from a genetic sub-isolate. This family segregates BD in at least four generations with 24 affected individuals.

Methods
We selected 16 family members for targeted sequencing. The selected individuals either carried the disease haplotype, were non-carriers of the disease haplotype, or served as married-in controls. We designed hybrid capture probes enriching for 5-9Kb fragments spanning the entire 12Mb region that were then sequenced to screen for candidate structural variants (SVs) that could explain the increased risk for BD in this extended family.

Results
Altogether, 201 variants were detected in the critically linked region. Although most of these represented common variants, three variants emerged that showed near-perfect segregation among all BD type I affected individuals. Two of the SVs were identified in or near genes belonging to the RNA Binding Motif Protein, X-linked (RBMX) gene family—a 330bp Alu (subfamily AluYa5) deletion in intron 3 of the RBMX2 gene and an intergenic 27bp tandem repeat deletion between the RBMX and G protein-coupled receptor 101 (GPR101) genes. The third SV was a 50bp tandem repeat insertion in intron 1 of the Coagulation Factor IX (F9) gene.
Conclusions

Among the three genetically linked SVs, additional evidence supported the *Alu* element deletion in *RBMX2* as the leading candidate for contributing directly to the disease development of BD type I in this extended family.

Introduction

Bipolar Disorder (BD) is a severe psychiatric disorder that is characterized by recurrent episodes of mania and depression. The lifetime prevalence is 2.4% [1] and the average age of onset is in the early twenties [1]. Based on twin studies the overall heritability for BD is 40–70%, and lifetime risk in first-degree relatives is 5–10% which is almost seven times higher than the general population risk [1].

Common complex disorders, such as bipolar disorder, are thought to be caused by a combination of genetic and environmental factors. To unravel the complex genetic makeup of these disorders, genome-wide association (GWAS) and family-based linkage studies have been applied. For BD, a number of significant linkage and association findings have been reported. One of these studies was by Pekkarinen et al. who reported significant linkage (lod score: 3.54) in a genome-wide survey to chromosome Xq24-q27 in a Finnish extended pedigree segregating BD in four generations [2]. A follow-up study based on microsatellite markers narrowed down the critically linked region to 12Mb [3]. Despite continued SNP mapping efforts, the critical region could not be further reduced, nor could a disease variant be pinpointed. This is not an uncommon phenomenon for common complex disease. To date, thousands of single nucleotide polymorphisms (SNPs) have been associated with various common complex disease phenotypes [4]; however, the effect sizes typically are small and explain only a portion of the disease heritability [5, 6]. Similar conclusions were drawn for the largest GWAS meta-analysis that was performed by the PGC Bipolar Disorder Working Group. They analyzed 20,353 cases and 31,358 controls, and identified 30 genome-wide significant loci, of which 20 were novel [7]. One emerging hypothesis holds that the missing heritability is hidden in other types of genetic variation than SNPs, such as structural variants (SVs) of >50 bp [8] in length, whose effects are not adequately represented by neighboring SNPs [5].

In the human genome there are over 25,000 SVs and, despite not being as common as SNPs, due to their size they make up 60% of all variant bases in the human genome [9]. Genome-oriented studies have shown that SVs like insertions, deletions, duplications, translocations, inversions, and tandem repeat expansions can all cause disease [10–12] for example through the disruption of key gene regulatory elements [5]. Over the past several decades, the discovery of new SVs has followed closely behind technological advances in sequencing platforms. Therefore, we wanted to explore the possibility of a SV being the driver of the linkage signal to chromosome Xq24-q27 in the extended pedigree enriched with BD originally reported by Pekkarinen and colleagues [3]. Altogether, 24 out of 61 (39.3%) members in this family were diagnosed with some form of psychiatric disorder (Fig 1), therefore presenting a much higher life-time prevalence than the general population (2.4%). The hypothesis was that this extended pedigree would present a less heterogenous form of bipolar disorder because it stemmed from a genetic sub-isolate in Finland and showed no significant linkage to other genomic regions. To this effect, we used long-read sequencing to screen the 12Mb genomic region for a potential disease-causing SV in key individuals for the extended pedigree. Single Molecule, Real-Time (SMRT) long-read sequencing (Pacific Biosciences, CA, USA) has shown

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**Competing interests:** JME, WJR and SK were full-time employees and shareholders of Pacific Biosciences. DR, GM and DLB were full-time employees of Roche Sequencing Solutions. The sequencing reagents and technology used is sold by Pacific Biosciences. The targeted enrichment kit used is sold by Roche Sequencing Solutions. The commercial affiliation does not alter our adherence to all PLOS ONE policies on sharing data and materials. All software, algorithms, protocols and methods used in this publication are publicly available. The samples and the data are legally owned by the THL Biobank, Finland and is not in the possession of the authors. The data contains sensitive information and can therefore not be shared publicly. Therefore, a request to access the data can be submitted to the ethics committee at https://thl.fi/en/web/thl-biobank/for-researchers/sample-collections.
to discover 80% more SVs than other sequencing methods [13] through recent human genome assembly studies. In addition, long sequencing reads also help resolve structural breakpoints and to define allele-specific haplotypes [14].

In this study we detected altogether 201 SVs from 16 key individuals in the critically linked 12Mb genomic region. Although most of these represented common variants that could be seen across many of the family members regardless of disease status, or in the general population, three of the SVs showed near-perfect segregation among affected individuals that were identified as carriers of the disease haplotype in the previous linkage studies [2, 3]. Two of these were located within the same gene family— one SV was a 330bp antisense Alu deletion in intron 3 of the RNA Binding Motif Protein, X-Linked 2 (RBMX2) gene, while another variant, a 27bp tandem repeat deletion, was located in the intergenic region of the RNA Binding Motif Protein, X-Linked (RBMX) and G protein-coupled receptor 101 (GPR101) genes. The third SV was a 50bp tandem repeat insertion located in intron 1 of the Coagulation Factor IX (F9) gene.

**Material and methods**

**Collection of study samples**

The extended family P101 was described in great detail in the original linkage studies [2, 3, 15]. The family originates from a genetic isolate in Eastern Finland where the population only stabilized in the 17th century, prior to a major expansion [16]. This was also one of the genetic hubs for the so-called Finnish disease heritage, which refers to a group of rare recessive disorders that due to genetic bottlenecks have become more prevalent in Finland than elsewhere in the world [17]. Pekkarinen et al. also alluded to BD being more enriched in this region compared to the nation in general [2].

Altogether, 24 out of 61 family members were diagnosed with a psychiatric disorder using the DSM-III-R (Diagnostic and Statistical Manual of Mental Disorder III–R) criteria. Out of these, 10 were diagnosed with BD (BD type I:8, BD type 2:1, BD not otherwise specified:1), one with schizoaffective disorder of bipolar type, five with recurrent major depression, two with schizophrenia, one with schizophreniform disorder, two with psychosis not otherwise
specified, and three with alcohol abuse. The clinical DSM-IV diagnosis for this pedigree was last updated in 2002, when all the individuals (except two: 686 was 33 years-of-age, and 690 was 24 years-of-age) were aged 42 years or older, well beyond the mean age of onset for bipolar disorder which is 22 years.

The study to uncover the genetic etiology of bipolar disorder in family P101 was approved by the Helsinki university hospital’s epidemiology and public health ethics committee on June 11th, 2003 (Dnro189/E3/2003). Participants gave a written informed consent.

We selected 16 individuals from this X-chromosomally linked family for targeted SMRT sequencing, including carriers of the disease haplotype, non-carriers of the disease haplotype, and non-related controls that were married into the family. All DNA samples sequenced represented the third and fourth generation of the extended family and included five cases of bipolar type 1 and two cases of recurrent major depression cases (Fig 1).

**Targeted enrichment and sequencing**

In order to survey the 12Mb significantly linked region, we used Roche Sequencing Solutions’ SeqCap EZ sequence capture technology in combination with SMRT® sequencing. A custom designed SeqCap EZ probe pool (Hoffman-La Roche, Basel, Switzerland) was built tiling nearly 77% of the 12Mb region (GRCh38 ChrX:128,049,971–139,850,027). The capture experiment was performed in accordance with the protocol PN 100-893-500-0, available from Pacific Biosciences. Specifically, 2μg of genomic DNA from each of the 16 samples were sheared to an average of 10Kb fragments using g-Tube™ by Covaris (PN 520079, Covaris, MA, USA). For each DNA sample, a library was made using the KAPA HyperPrep Kit® for Illumina (Hoffman-La Roche). Libraries were amplified (12 cycles) under conditions optimized for long fragments before size selection using the BluePippin® System (Sage Science, MA, USA) to a desired size ranged of 5-9Kb. Next, the SeqCap EZ probes were hybridized to 1.5μg of each DNA library, followed by bead capture and washing. The captured DNA fragments were then amplified (18 cycles) and PacBio SMRTbell® sequencing libraries were prepared as recommended by the manufacturer. The final libraries were size selected on the BluePippin to a size of 6–6.5Kb. All size determinations were performed using a Fragment Analyzer (Agilent Technologies, CA, USA). The samples were not multiplexed during the library preparation process and were sequenced separately on individual SMRT Cells (3 for each sample, except for one sample with 2 SMRT Cells) on the PacBio RS II instrument (Fig 2).

**Sequencing analysis and visualization of results using Integrated Genome Viewer (IGV)**

PacBio continuous long reads were mapped to the GRCh38 reference genome using pbmm2 v0.12.0 (https://github.com/PacificBiosciences/pbmm2), using the ‘—median-filter’ flag to align one subread per Zero-Mode Waveguide (ZMW). Putative PCR duplicates were flagged with a custom script (https://github.com/williamrowell/markdup). The on-target rate was calculated as the proportion of ZMWs with a read overlapping the targeted region after deduplication. The results were visualized using IGV 2.5 [18] with Quick Consensus mode enabled and indels shorter than 3bp hidden.

**Structural variant detection**

SVs were detected in the deduplicated alignments with the pbsv 2.0.1 workflow (https://github.com/PacificBiosciences/pbsv). The pbsv software will detect variants larger than 20bp. Using bedtools v2.27.1, we filtered for variants in the targeted region. Using bcftools v1.9, we filtered for variants with the requirement that an SV must be detected at least once in BD-
affected individuals (685, 691, 698, 699, 700) and absent in the outgroup of non-X-chromosomal relatives (689, 690, 693, 697).

Verification of the 330bp \textit{Alu} element deletion using PCR and Sanger Sequencing

Forward and reverse primers (RBMX2\textsubscript{F}: 5\textquotesingle -ACATTGCCAAATTGCTCTCC-3\textquotesingle and RBMX2\textsubscript{R}: 5\textquotesingle -CACCACCACACCTGGCTAC-3\textquotesingle) were designed to amplify the flanking region of \textit{Alu} deletion in the intron 3 region of the \textit{RBMX2} gene. To be able to amplify both the reference allele and the deletion allele within \textit{RBMX2} (S1 Table), we designed two additional primers (RBMX2\textsubscript{Rref}: 5\textquotesingle -CATATCTGACACCTTTAATTTCTA-3\textquotesingle and RBMX2\textsubscript{Rdel}: 5\textquotesingle -CATATCTGACACCTTTAATTTCTG). Reverse primers (RBMX2\textsubscript{Rref} and RBMX2\textsubscript{Rdel}) have a single nucleotide difference at the last 3\textquotesingle position of the primer (A/G, shown in bold type here). This position is ChrX:130,406,896 in GRCh38 and also a known SNP (rs56113207). Two allele specific PCR reactions (RBMX2\textsubscript{F} and RBMX2\textsubscript{Rref} (1215bp, reference allele); RBMX2\textsubscript{F} and RBMX2\textsubscript{Rdel} (896bp, deletion allele) were performed for each of the 16 individuals.

All PCR reactions were performed in 50\textmu l total reaction volume for each sample: DNA 1\textmu l, 10x Buffer B1 5\textmu l, MgCl\textsubscript{2} (25mM) 3\textmu l; dNTPs (10mM) 1\textmu l; Primers (10\mu M) 2.5\textmu l each; Phusion\textsuperscript{R} Hot Start II DNA Polymerase (2U/\mu l, ThermoFisher Scientific) or Hot FirePol Polymerase (5U/\mu l, Solis BioDyne, EE) 0.5\textmu l; H\textsubscript{2}O 34.5\textmu l. PCR reactions were performed on Veriti 96 well thermal cycler (ThermoFisher Scientific). Cycling conditions were as follows: polymerase activation 95\degree C for 15sec, then 35 amplification cycles of 95\degree C for 15sec, 63\degree C for 30sec and 72\degree C for 60sec with a final extension 72\degree C for 5min. PCR products were purified with AMPure\textsuperscript{R} Beads (Beckman Coulter, CA, USA).

The PCR product of RBMX2\textsubscript{F} and RBMX2\textsubscript{Rdel} (893bp) was sequenced using forward and reverse primers and Sanger sequencing. The longer reference-like i.e. sequence equivalent
to current human genome sequence (Hg38), PCR product (1215bp) was sequenced using the forward and reverse primer (RBMX2_RRef). To resolve the obtained hairpin structure the longer reference-like PCR product was first cut with Scal restriction enzyme and then sequenced with forward (RBMX2_F) and reverse (RBMX2_RRef) and two additional primers BRM X2_389 (5’ –TCGATCTTTGACCTGTA-3’) and BRMX2_397 (5’ –CGGATCACGAG GTCAAGAGA-3’). Finally, the original longer PCR product was cut with BbvI followed by sequencing with RBMX2_Long_BbvI_F (5’ –TCATATCCTTTGCCAACTTTC-3’) primer.

All Sanger sequencing used BigDye chemistry and was performed on Applied Biosystems 3130 Genetic Analyzer (ThermoFisher Scientific) for a small subset of family members (693, 688 and 700) that could serve as representatives for the other samples that showed the same variations based on PCR results and under the assumption that they would generate same Sanger sequencing results. To predict secondary structures of PCR amplified sequences we used Secondary Structure Web Server (http://rna.urmc.rochester.edu/RNAstructureWeb/Servers/Predict1/Predict1.html) with default structure options, selecting DNA was as a nucleic acid type.

Sequence features and genome conservation analysis
Dfam (release 3.1, dfam.org/home) database was used to identify the family of Alu elements within intron 3 of RBMX2 gene. Sequence homologies of the highlighted Alu element in intron 3 and the surrounding intron 3 sequence was explored using default parameters with the LAST [19] alignment tool and blastn [20]. In order to visualize genome conservation between species for the 330bp Alu element deletion in the RBMX2 gene, the comparative genomics track in www.ensemble.org was utilized.

Results
Targeted enrichment using customized SeqCap EZ probe panel
SeqCap EZ capture efficiency ranged from 51% to 69%, with a mean of 61% on-target reads (S2 Table). Altogether 47 SMRT Cells were sequenced, with mean throughput of 572Mb per SMRT Cell. Unique molecular coverage of the targeted region ranged from 14.6-fold to 34.5-fold, with a mean of 21-fold sequencing coverage (Fig 3).

Structural variation detection analysis
Using the PBSV workflow, we identified 201 SVs within the 12Mb target region, however most represented common variants that could be seen across many of the family members, regardless of the disease status (S3 Table). From the set of variants seen in known carriers of the disease haplotype (685, 691, 698, 699, 700), we subtracted any variant seen in the haplotype non-carriers or married-in controls (689, 690, 693, 697), identifying 35 variants unique to carriers. Of these, only three variants were well-supported in four or more known carriers; a 330bp antisense Alu deletion in intron 3 of the RBMX2 gene (ChrX:130,405,824–130,406,154), a 27bp tandem repeat deletion in the intergenic region between the RBMX and GPR101 genes (ChrX:137,012,126–137,012,153) and a 50bp tandem repeat insertion in intron 1 of the F9 gene (ChrX:139,536,239) (Fig 4).

Verification of the RBMX2 330bp Alu element deletion using PCR and Sanger sequencing
We designed one forward and two allele specific reverse primers to amplify both reference-like and deletion alleles in region of RBMX2, intron 3. Based on PacBio reads, all 5 BPD-diagnosed
individuals and four non-diagnosed but X-chromosomally linked individuals (687, 688, 694, 696) carried the C allele of this SNP while all others carried the T allele. Allele frequencies for this SNP in all populations tested are T: 84.3% and C: 15.7% and for the Finnish population T: 87.5% and C:12.5%, respectively (www.ensemble.org).

We used PCR and Sanger sequencing to verify the RBMX2 gene 330bp deletion. One forward and two allele specific reverse primers were designed to amplify the reference like region and the flanking region of the 330bp deletion (Fig 5A). DNA from 16 family members were PCR amplified with allele specific primer pairs RBMX2_F and RBMX2_Rref, RBMX2_F and RBMX2_Rdel (Fig 5B). Two different DNA polymerases, Phusion Hot Start II DNA Polymerase and HOT FIREPol™ DNA Polymerase, were tested. We observed that HOT FIREPol DNA Polymerase performed better, presumably due to a lack of 3’ → 5’ exonuclease activity.

Sanger sequencing was performed for a subset of the P101 family members (693, 688, and 700). The deletion allele (Fig 5B, wells 4 and 6) was sequenced using the RBMX2_F and RBMX2_Rdel primers. On our first attempt, we failed to sequence the longer PCR product of the reference allele (Fig 5B, wells 1 and 3). We hypothesize that the local sequence context generated a secondary hairpin structure due to consecutive opposite strand Alu elements after the denaturation step, therefore preventing elongation during Sanger sequencing. Subsequently, we used Scal and BbvI restriction enzymes to cut the PCR product and disrupt hairpin formation before sequencing.

Sequence features and genome conservation in the candidate Alu element

The highlighted RBMX2 Alu element deletion (ChrX:130,405,825–130,406,154) is on the antisense strand in relation to the RBMX2 gene and it contains all sequence features characteristic
to Alu elements [21] (Fig 6A). These consist of a nine bp (TGCTTTGCC) direct repeat and target site duplications (TDSs) that flank the Alu element. Also, in the middle, the Alu element contains a A₅TACA₅-box as well as a 39 bp long A-tail on the 3'end (Fig 6B and 6C). After the 330bp deletion only one direct repeat remains of the Alu sequence. Based of Dfam database search, the Alu sequence is identified as AluYa5 subfamily (e-value 2.2e-115). In order to explore how the 330bp Alu deletion may affect secondary structures several different sequence alignments were performed. First, the RBMX2 gene (11.76Kb) and intron 3 (4.81Kb) sequences were aligned, both with and without the Alu deletion against itself (Fig 7A–7C). Most of the homologies seen are short but represent almost the full length of the Alu element like sequences.

 Altogether, four Alu elements, identified using Dfam, two (AluSx and AluSx1) on the sense and two (AluYa5 and AluJr) on the antisense strand, are found in intron 3 of the RBMX2

| SV # | SV type                  | Size  | Location                        | Carriers of SV                  | Verification methods                  |
|------|--------------------------|-------|---------------------------------|---------------------------------|---------------------------------------|
| 1.   | Alu deletion             | 330 bp| ChrX:130,405,824 RBMX2 intron 3| 685, 687, 688, 691, 694, 696, 698, 699, 700 | PBSV workflow, PCR+Sanger, manual data inspection |
| 2.   | Tandem repeat deletion   | 27 bp | ChrX:137,012,126 intergenic between genes RBMX and GPR101 | 685, 686, 687, 688, 691, 694, 695, 696, 698, 699, 700 | PBSV workflow |
| 3.   | Tandem repeat insertion  | 50 bp | ChrX:139,536,239 F9 intron 1   | 685, 687, 688, 691, 692, 694, 695, 696, 698, 699, 700 | PBSV workflow |

Fig 4. Candidate SVs. In total, three SVs were highlighted in this study. These included an Alu deletion (1, red), a tandem repeat deletion (2, blue) and a tandem repeat insertion (3, black) of various sizes. Two of them (1 and 2) were located in or in close vicinity of different RBMX gene family members, while one was located in the F9 gene. The pedigree shows the segregation by number of each of the SVs.

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gene (see Table 1). The consecutive opposite Alu elements \((\text{AluSx}, \text{AluYa5}, \text{AluSx1} \text{and} \text{AluJr})\) with reference-like sequences would be expected to form hairpin loops in pre-mRNA as shown in Fig 8A, however at the DNA level, given the close proximity of the Alu elements, these may form a G quadruplex structure (Fig 8B). The homology between the four Alu elements can clearly be seen in Fig 8C, when comparing the sequence of the three reference-like Alu elements in intron 3 to the highlighted Alu element.

To further explore the potential structural and functional relevance of this Alu element, the Alu sequence and flanking region was compared against the reference genomes of other primates, including chimpanzee, gorilla, macaque and orangutan. The 330bp Alu deletion allele was not observed in any of these non-human primates (Fig 9).

**Discussion**

An extended pedigree segregating BD in four generations was originally found to be linked (lod score: 3.54) to Xq24-q27 in a genome-wide survey using microsatellite markers [2]. A follow-up study narrowed down the critically linked region to 12Mb [3]. In order to pinpoint disease associated variants, we used long-read sequencing to fully characterize the 12Mb genomic region in 16 out of the 61 family members. The P101 pedigree, which originates from a genetic isolate in Northern Finland, portrays a highly elevated recurrence risk for BD compared to the general population [1]. It has been hypothesized that the genetic heterogeneity would be reduced in isolated populations, even for genetically complex disorders such as BD [22]. This seems to also be the case for this extended family as it showed no evidence of significant linkage to any other chromosomal region in the original genome-wide scans [3], therefore this family may display a more monogenic form of the disorder.
In order to survey the critically linked 12Mb region, Roche’s hybridization-based target enrichment in combination with PacBio long-read sequencing was used. The SeqCap EZ probe pool was designed to enrich for 5-9Kb fragments spanning the entire 12Mb region. Compared to more traditional sequencing technologies, SMRT Sequencing has previously shown to be more efficient in capturing the full range of variations, including larger structural variations [10]. This technology has also demonstrated the ability to overcome other short-
Table 1. *Alu* elements in the RBMX2 gene.

| Sequence name | Model accession | Model name | Bit score | e-value | Model start | Model end | Strand | Alignment start | Alignment end | Envelope start | Envelope end |
|---------------|-----------------|------------|-----------|---------|-------------|-----------|--------|----------------|---------------|---------------|-------------|
| Intron3       | DF0000047       | AluSx      | 340.6     | 1.6e-101| 1           | 308       | +      | 647            | 957           | 647           | 960         |
| Intron3       | DF0000303       | L1ME4a_3end| 30.8      | 1.1e-07  | 725         | 869       | -      | 1084           | 963           | 1096          | 962         |
| Intron3       | DF0000303       | L1ME4a_3end| 118.0     | 4.7e-34  | 160         | 504       | -      | 1390           | 1099          | 1409          | 1088        |
| Intron3       | DF0000053       | AluYa5     | 382.8     | 3.1e-114 | 1           | 310       | -      | 1704           | 1394          | 1704          | 1374        |
| Intron3       | DF00000303      | L1ME4a_3end| 69.0      | 3.1e-19  | 1           | 172       | -      | 1877           | 1708          | 1877          | 1705        |
| Intron3       | DF0000048       | AluSx1     | 320.7     | 1.8e-95  | 1           | 306       | +      | 1942           | 2246          | 1942          | 2250        |
| Intron3       | DF0000845       | Tigger4a   | 28.8      | 2.2e-06  | 199         | 236       | -      | 1947           | 1910          | 1954          | 1910        |
| Intron3       | DF0000846       | Tigger4b   | 241.9     | 2.1e-71  | 1           | 341       | -      | 2565           | 2243          | 2565          | 2240        |
| Intron3       | DF0000281       | L1MC5a_3end| 24.2      | 1.3e-06  | 1566        | 1694      | -      | 2688           | 2567          | 2690          | 2546        |
| Intron3       | DF0000010       | L1MC4_3end | 46.4      | 2.2e-13  | 2679        | 2786      | -      | 2812           | 2703          | 2834          | 2691        |
| Intron3       | DF0000147       | FRAM       | 67.4      | 3.00E-18 | 66          | 168       | -      | 2901           | 2799          | 2917          | 2791        |
| Intron3       | DF0000008       | L1M5_orf2  | 20.0      | 1.5e-05  | 1915        | 2111      | -      | 3061           | 2909          | 3087          | 2900        |
| Intron3       | DF0000035       | AluJr      | 247.8     | 2.5e-73  | 4           | 311       | -      | 3506           | 3200          | 3509          | 3195        |
| Intron3       | DF00001071      | SVA_E      | 84.0      | 2.3e-24  | 133         | 317       | +      | 3626           | 3835          | 3625          | 3855        |
| Intron3       | DF0000144       | FLAM_C     | 106.5     | 3.2e-30  | 1           | 139       | -      | 3680           | 3552          | 3680          | 3533        |
| Intron3       | DF0000144       | FLAM_C     | 122.7     | 3.3e-35  | 1           | 139       | -      | 3847           | 3709          | 3847          | 3689        |
| Intron3       | DF0000008       | L1M5_orf2  | 120.3     | 9.1e-36  | 63          | 1042      | -      | 4773           | 3848          | 4794          | 3846        |

The four different types of *Alu* elements (bolded) were identified in intron 3 of the RBMX2 gene. Two were found on the sense strand and two on the antisense strand. Identification was done using Dfam.

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**Fig 8. A schematic of the secondary structure of intron 3 in the RBMX2 gene.** Altogether four *Alu* elements can be found within this intron. The gray boxes represent reference like *Alu* elements while the blue *Alu* box represents the *Alu* element with the 330bp deletion. The sense and antisense strands are marked with a plus and minus sign, respectively. (a) The pre-mRNA secondary loops are shown for the *Alu* elements of opposite orientation. (b) A DNA *Alu* G-quadruplex structure is shown as described by Larsen et al 2018(22). (c) The full extent of the homologies between the *Alu* elements and the RBMX2 gene intron 3 sequence are shown in this dotplot. The black arrows indicate the start and end positions of the *Alu* element with the 330bp deletion.

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read sequencing barriers like accessing difficult to sequence regions such as long repeat expansions [23] with uniform and unbiased sequencing coverage [24]. The multi-kilobase sequence reads also help allelic phasing across large regions [25].

The SV analysis identified in total 201 SVs among the sixteen individuals in the 12Mb genomic region, however, the majority represented common variants seen across many family members regardless of disease status and therefore are not likely to be linked to BD. Nevertheless, three SVs emerged that showed verifiable segregation among the affected family members only.

Interestingly, two of the SVs were found in or near the RBMX-gene family; a 330bp antisense Alu element deletion in the intron 3 of the RBMX2 gene and a 27bp tandem repeat deletion in the intergenic region between the RBMX and GPR101 genes. The third SV was a 50bp tandem repeat insertion in the F9 gene, which is a coagulation factor associated with HEMB (Hemophilia B) and THPH8 (Thrombophilia, X-Linked, due to factor IX defect). Although not an obvious candidate gene for the etiology of BD, the F9 gene does encode for vitamin K-dependent coagulation Factor IX that circulates in the blood, and alterations of this gene, including point mutations, insertions and deletions, cause Factor IX deficiency. This is a recessive X-linked disorder, also called hemophilia B or Christmas disease that has shown linkage to both major depressive disorder as well as bipolar disorder in past studies [26–28].

The RBMX and RBMX2 genes, located on Xq26, encode RNA-binding proteins that have several roles in the regulation of pre- and post-transcriptional processes [29, 30]. They have been implicated in tissue-specific regulation of gene transcription and alternative splicing of several pre-mRNAs that can either activate or suppress exon inclusion [29, 30]. Mutations in RBMX have also been associated with X-linked syndromic mental retardation-11(MRXS11; OMIM 300238). Shashi and colleagues identified a hemizygous 23bp deletion in exon 9 of the RBMX gene, resulting in a frameshift and premature termination [31, 32]. The mutation, which was found by whole-exome sequencing and confirmed by Sanger sequencing, segregated with the disorder in the family and was not found in the Exome Sequencing Project database or in 1300 controls. Here, we detected a 27bp tandem repeat deletion in the intergenic region between RBMX and the GPR101 genes that segregated together in all the BD type I diagnosed P101 family members, as well as in one affected with major depression. In addition, four unaffected family members (687, 688, 694, 696) were also carriers of the tandem repeat variant. The GPR101 gene is primarily expressed in brain and encodes a G protein-coupled receptor. G protein-coupled receptors are embedded in the outer membrane of cells, where they relay chemical signals from outside of the cell to the interior. The GPR101 protein is predominantly expressed in the pituitary gland during fetal development and again at adolescence, two stages noted for rapid body growth [33, 34].
The RBMX2 gene, located on Xq26.1 is over 11Kb in length and involved in pre-mRNA splicing. Here, we found a 330bp antisense Alu element deletion in intron 3 that again segregated in all P101 family members affected with BD type I, in addition to the same four unaffected family members (687, 688, 694, 696) as in the RBMX gene. All but 696 are females that might exhibit non-random X-inactivation where more of the mutant allele is expressed than in the unaffected female carriers in the family. The unaffected male 696 was 43 years old when initially entered into the study in 2002 and might have been pre-symptomatic with an expected later age of onset or simply a case of non-penetration. The fact that the mean age of onset in this extended pedigree was 21 years would suggest the latter.

Like the RBMX gene, RMBX2 has been shown to play a central role in brain development and function. ClinVar reported (submission accession #SCV000239951.1) on a SNP (rs5977266) in exon 6 that was characterized as a missense mutation and protein change (R287H) involved in abnormal neuronal migration. This finding falls within a diverse group of congenital brain developmental disorders that are characterized by defects in neuronal migration in the brain during early fetal development. These neuronal migration defects can result in brain abnormalities that are usually manifested with mental retardation and epilepsy [35]. The RBMX2 gene is also located adjacent to the Solute Carrier Family 25 Member 14 (SLC25A14) gene, which has been shown to have altered expression in autism patients [36].

The highlighted 330bp antisense Alu element in the intron 3 of the RBMX2 gene contained a direct repeat (TGCTTTGCC) flanking the element, a relatively long polypyrimidine tract, and a A5TACA5 box. Alu elements are known to be one of the most abundant repetitive elements in the human genome, with >1.3 million copies accounting cumulatively for at least 11% of human genomic DNA [22, 37]. They are approximately 300 nucleotides in length and tend to be accumulated in GC-rich regions that participate in the architecture of the genome by delimiting the active/inactive domains and the epigenetic landscape and gene regulation at different levels [38].

Alu elements are a highly successful family of primate-specific retrotransposons that have fundamentally shaped primate evolution, including human evolution [39]. The Alus play critical roles in the formation of neurological networks and the epigenetic regulation of biochemical processes throughout the central nervous system, and thus are hypothesized to have contributed to the origin of human cognition [39]. Despite the benefits that Alu elements may provide, deleterious Alu activity is associated with at least 37 neurological and neurodegenerative disorders, wherein Alu elements are hypothesized to disrupt key cellular processes that result in or contribute to the disease state [39]. Alu elements have many modes of action ranging from novel gene formation, elevated transcriptional diversity, long non-coding RNA and microRNA evolution (including circular RNAs), transcriptional regulation, and creation of novel response elements [40–44]. Moreover, Alu elements fundamentally alter the three-dimensional architecture and spatial organization of primate genomes by defining the boundaries of chromatin interaction domains (i.e., topologically associating domains (TADs)) [45]. The deleted Alu element (subfamily AluYa5) in our current study is located on the antisense strand in relation to RBMX2 gene. The AluY lineage is the youngest of the Alu lineages and has the largest number of functionally active sequences [46]. We observed that in the current human reference sequence (Hg38) the rather long (39 bp) stretch of homopolymeric adenosines (A-tail) we observed at the 3' end of the Alu element here could indicate a more active Alu element [47].

It is known that polypyrimidine tracts within pre-messenger RNA (mRNA) promote the assembly of the spliceosome, the protein complex specialized for carrying out RNA splicing during the process of post-transcriptional modification [48]. Therefore, a deletion of a polypyrimidine tract may contribute to a change in 3' splice site recognition. Also, the absence of
polypyrimidine tract in pre-mRNA might have an impact on the polypyrimidine tract binding protein 1 (PTB1) involved in RNA looping when two separate pyrimidine tracts are present within the same RNA [49]. In fact, there are examples where mutations that alter polypyrimidine tract sequences may disrupt pre-mRNA splicing, leading to exon skipping or shortening, partial or full intron retention in the mRNA [50]. There is no specific known mechanism for removal of a complete Alu element [51], but the direct identical repeats flanking the Alu element are believed to be involved in recombination [50]. Here we found two direct repeats of 9 bp long (TGCTTTGCC) TSDs sequences flanking the intact Alu element allele whereas only one remained at the deletion allele. In addition to the complete Alu element in intron 3 of the RBMX2 gene, we found three additional young Alu elements (AluSx, AluSx1 and AluJr) in the intron 3 region. The consecutive elements were in close proximity on opposite strands, which opens up the possibility of them forming pre-mRNA secondary loops or on the DNA-level, Alu G-quadruplex structures [39]. These non-B DNA Alu G-quadruplex structures are shown to serve as potential binding sites for proteins such p53, tumor suppressor protein, involved in complicated regulatory network by activating and repressing expression of ~1000 human genes [52]. In addition, Cui et al observed that the strongest binding sites for p53 reside in the relatively young Alu elements located in introns [52].

The RBMX2 gene in itself is an important protein for the first step of splicing [53] which could further compromise transcription of candidate genes if compromised. In addition, we also found no evidence of the presence of this Alu element in other primate species, therefore making it specific to humans which would be expected if it was part of the mechanism leading to psychiatric disease.

In conclusion, targeted long read sequencing using probe-based hybrid-capture serves as a powerful tool to characterize the structure of regions linked to or associated with common complex disorders and may ultimately help pinpoint functional variants underlying the developmental or biochemical etiology of those disorders. In this study three SVs were highlighted, each an interesting candidate as a disease-causing variant for BD. However, the 330bp Alu deletion in the RBMX2 gene emerged as the strongest candidate due to evidence supporting the hypothesis that loss of a complete Alu (subfamily AluYa5) element from within the RBMX2 gene may be involved in the disease development of BD type I in this extended family. Here, we located two opposite strand Alu-Alu element pairs in intron 3 of this gene. This type of consecutive sense and antisense Alu element can potentially at pre-mRNA level form long double stranded RNA (dsRNA) structures and play a role in regulation of alternative splicing [54]. Due to this antisense 330bp Alu deletion, one of the Alu-Alu base-pairings cannot form the dsRNA, therefore this Alu deletion may influence the RNA splicing and generate novel transcripts in the RBMX2 gene. In order to fully understand the exact mechanism of how this 330bp deletion would be involved in the development of BD type I—further studies are warranted. For example, full-length isoform characterization and expression studies of the RBMX2 gene would allow better delineation on how the downstream gene products are affected by this variant and how it could contribute to disease development. Additional, functional studies may also be warranted such as mouse models to fully elucidate the underlying disease mechanism.

**Supporting information**

**S1 Raw images. Raw blot/gel image.** The raw image for Fig 5.
(PDF)

**S1 Table. PCR experiment details.** A description of wells, sample identification numbers and primer pairs used in the PCRs illustrated in the gel picture (Fig 5B)
(DOCX)
S2 Table. Sequencing on-target rates. Sequencing summary for 16 family members. On-target rate is calculated as the number of unique templates for a given sample after deduplication. (DOCX)

S3 Table. Sequencing genotypes. Genotypes for all 16 family members at positions of the 35 variants identified as present in carriers and absent in non-blood related controls. Positions are provided in GRCh38 coordinates. In variant length column, positive sizes indicate insertions, and negative sizes indicate deletions. Genotypes are reported by PBSV as diploid, and encoded as 0 (reference allele), 1 (alternate allele), or (undetermined). Homozygous reference (0/0) or undetermined phenotypes (,/) are shown in black, and homozygous alternate (1/1) or heterozygous (0/1) phenotypes are shown in red. Below the genotypes the number of supporting reads (REF, ALT) are shown in parentheses the total informative reads used by PBSV (which may not equal total overall coverage) are show in square brackets. The three highlighted variants are bolded and marked with an asterisk. The P101 family member affected with BD type I are underlined. (DOCX)

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