

1 **Comprehensive cross-disorder analyses of *CNTNAP2* suggest it is unlikely to be a**
2 **primary risk gene for psychiatric disorders.**

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19 **Short title:** Role of *CNTNAP2* gene in psychiatry

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25 **Word limits for PLoS Genetics (Archival Report):**

26

27 Abstract (max=300 words): 283

28 Article body: 3,950

29 Tables: 5 Tables

30 Figures: 2 Figures

31 Supplemental information: 1 document

32

33 **ABSTRACT**

34 The contactin-associated protein-like 2 (*CNTNAP2*) gene is a member of the neurexin
35 superfamily. *CNTNAP2* was implicated in the cortical dysplasia-focal epilepsy (CDFE)
36 syndrome, a recessive disease characterized by intellectual disability, epilepsy, language
37 impairments and autistic features. Associated SNPs and heterozygous deletions in *CNTNAP2*
38 have also frequently been reported in autism, schizophrenia and other psychiatric or
39 neurological disorders. We aim here to gain conclusive evidence for the role of *CNTNAP2* in
40 susceptibility to psychiatric disorders by the comprehensive analysis of large genomic datasets.
41 In this study we used: i) summary statistics from the Psychiatric Genomics Consortium (PGC)
42 GWAS; ii) examined all reported *CNTNAP2* structural variants in patients and controls; iii)
43 performed cross-disorder analysis of functional or previously associated SNPs; iv) and
44 conducted burden tests for pathogenic rare variants using sequencing data (4,483 ASD and
45 6,135 schizophrenia cases, and 13,042 controls).

46 In a CNV microarray study, we previously identified a 131kb deletion in *CNTNAP2* intron 1,
47 removing a FOXP2 transcription factor binding site in an extended BD family. Here we
48 perform a quantitative-PCR validation showing imperfect segregation with disease (5 bipolar
49 disorder relatives). The distribution of CNVs across *CNTNAP2* in psychiatric cases from
50 previous reports was no different from controls of the database of genomic variants. Gene-
51 based association testing did not implicate common variants in autism, schizophrenia or other
52 psychiatric phenotypes. The association of proposed functional SNPs rs7794745 and
53 rs2710102, reported to influence brain connectivity, was not replicated; nor did functional
54 SNPs yield significant results in meta-analysis across psychiatric disorders. Disrupting
55 *CNTNAP2* rare variant burden was not higher in autism or schizophrenia compared to controls.
56 This large comprehensive candidate gene study indicates that *CNTNAP2* may not be a robust
57 risk gene for psychiatric phenotypes.

58 **AUTHOR SUMMARY**

59 Genetic mutations that disrupt both copies of the *CNTNAP2* gene lead to severe disease,
60 characterized by profound intellectual disability, epilepsy, language difficulties and autistic
61 traits. Researchers hypothesized that this gene may also be involved in autism given some
62 overlapping clinical features with this disease. Indeed, several large DNA deletions affecting
63 one of the two copies of *CNTNAP2* were found in some patients with autism, and later also in
64 patients with schizophrenia, bipolar disorder, ADHD and epilepsy, suggesting that this gene
65 was involved in several psychiatric or neurologic diseases. Other studies considered genetic
66 sequence variations that are common in the general population, and suggested that two such
67 sequence variations in *CNTNAP2* predispose to psychiatric diseases by influencing the
68 functionality and connectivity of the brain. In the current study, we report the deletion of one
69 copy of *CNTNAP2* in a patient with bipolar disorder from an extended family where five
70 relatives were affected with this condition. To better understand the involvement of *CNTNAP2*
71 in risk of mental illness, we performed several genetic analyses using a series of large publically
72 available or in-house datasets, comprising many thousands of patients and controls. Despite
73 the previous consideration of *CNTNAP2* as a strong candidate gene for autism or schizophrenia,
74 we show that neither common, deletion nor ultra-rare variants in *CNTNAP2* are likely to play
75 a major role in risk of psychiatric diseases.

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83 INTRODUCTION

84 The contactin-associated protein-like 2 (*CNTNAP2*) is located on chromosome 7q35-36.1, and
85 consists of 24 exons spanning 2.3Mb, making it one of the largest protein coding genes in the
86 human genome. This gene encodes the CASPR2 protein, related to the neurexin superfamily,
87 which localises with potassium channels at the juxtaparanodal regions of the Ranvier nodes in
88 myelinated axons, playing a crucial role in the clustering of potassium channels required for
89 conduction of axon potentials [1]. *CNTNAP2* is expressed in the spinal cord, prefrontal and
90 frontal cortex, striatum, thalamus and amygdala; this pattern of expression is preserved
91 throughout the development and adulthood [2, 3]. Its function is related to neuronal migration,
92 dendritic arborisation and synaptic transmission [4]. The crucial role of *CNTNAP2* in the
93 human brain became clear when Strauss *et al*, reported homozygous mutations in Old Order
94 Amish families segregating with a severe Mendelian condition, described as cortical dysplasia-
95 focal epilepsy (CDFE) syndrome (OMIM 610042) [5]. In 2009, additional patients with
96 recessive mutations in *CNTNAP2* were reported, with clinical features resembling Pitt-Hopkins
97 syndrome [6]. To date 33 patients, mostly from consanguineous families, have been reported
98 with homozygous or compound deletions and truncating mutations in *CNTNAP2* [5-9], and are
99 collectively described as having CASPR2 deficiency disorder [7]. The common clinical
100 features in this phenotype include severe intellectual disability (ID), seizures with age of onset
101 at two years and concomitant speech impairments or language regression. The phenotype is
102 often accompanied by dysmorphic features, autistic traits, psychomotor delay and focal cortical
103 dysplasia.

104 *CNTNAP2* is also thought to contribute to the phenotype in patients with interstitial or
105 terminal deletions at 7q35 and 7q36. Interstitial or terminal deletions encompassing *CNTNAP2*
106 and several other genes have been described in individuals with ID, seizures, craniofacial
107 anomalies, including microcephaly, short stature and absence of language [10]. The severe

108 language impairments observed in patients with homozygous mutations or karyotypic
109 abnormalities involving *CNTNAP2* suggested a possible functional interaction with *FOXP2*, a
110 gene for which heterozygous mutations lead to a monogenic form of language disorder [11].
111 Interestingly, *Vernes et al.*, found that the FOXP2 transcription factor has a binding site in
112 intron 1 of *CNTNAP2*, regulating its expression [12]. Considering that a large proportion of
113 autistic patients show language impairments and most individuals with homozygous mutations
114 in *CNTNAP2* manifest autistic features, several studies investigated the potential involvement
115 of *CNTNAP2* in autism spectrum disorder (ASD). In particular, two pioneering studies showed
116 that single nucleotide polymorphism (SNP) markers rs2710102 and rs7794745 were associated
117 with risk of ASD [13, 14]. Moreover, in subsequent studies, rs2710102 was implicated in early
118 language acquisition in the general population [15], and showed functional effects on brain
119 activation in neuroimaging studies [16-19]. Furthermore, genotypes at rs7794745 were
120 associated with reduced grey matter volume in the left superior occipital gyrus in two
121 independent studies [20, 21], and alleles of this SNP were reported to affect voice-specific
122 brain function [22]. Genetic associations with ASD for these, and several other SNPs in
123 *CNTNAP2*, have been reported in a number of studies [23-28]. Along with the first reports of
124 variants associated with ASD, copy number variant (CNV) deletions have also been described
125 in ID or ASD patients, which were proposed to be highly penetrant disease-causative mutations
126 [13, 29-38]. To better understand the role of *CNTNAP2* in ASD pathophysiology, knockout
127 mice were generated. Studies of these mice reported several neuronal defects when both copies
128 of *CNTNAP2* are mutated: abnormal neuronal migration, reduction of GABAergic
129 interneurons, deficiency in excitatory neurotransmission, and the delay of myelination in the
130 neocortex [2, 39, 40].

131 These intriguing findings prompted additional investigations of *CNTNAP2* across other
132 psychiatric disorders or language-related traits, with additional reports of variants being

133 associated with schizophrenia (SCZ), bipolar disorder (BD), specific language impairment
134 (SLI) and several other phenotypes or traits [12, 15, 41-50]. Consecutively, other studies
135 reported CNV deletions in *CNTNAP2* in other psychiatric phenotypes such as schizophrenia
136 [51, 52], bipolar disorder [52-54], and ADHD [55]; neurological disorders [56-61]; and
137 language-related phenotypes [61-65]. Interestingly, several of these structural variants were
138 found in intron 1 of *CNTNAP2*, encompassing the FOXP2 transcription factor binding site.

139 Our group recently performed CNV analysis in extended families with bipolar disorder
140 [66], and found an intronic deletion in one individual which removed the FOXP2 binding site,
141 prompting the need for a segregation analysis in this family. We therefore aimed in this current
142 study to examine the evidence for a role of the *CNTNAP2* gene in multiple psychiatric
143 phenotypes, performing a comprehensive analysis of common and rare variants, CNVs and *de*
144 *novo* mutations using both in-house data and publically available datasets.

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146 RESULTS

147 *Examination of an intronic deletion in CNTNAP2 in an extended family with bipolar disorder*

148 CNV microarray analysis was performed in two affected individuals from an extended family
149 which included five relatives affected with bipolar I disorder. A drop in signal intensity for 340
150 consecutive probes was compatible with a deletion of 131 kb in intron 1 of *CNTNAP2*
151 (hg19/chr7:146203548-146334635; Fig 1A). The deletion encompasses the described binding
152 site for the transcription factor FOXP2 (hg19/chr7:146215016-146215040) [12]. The deletion
153 was detected in one of the two affected individuals examined. To infer deletion segregation
154 amongst relatives, WES-derived genotypes were used to create haplotypes across chromosome
155 7q35 (Fig 1B). The WES-derived haplotype analysis was uninformative due to incomplete
156 genotype data (unaffected descendants of deceased patient 8404 not included in the WES
157 study) and a likely recombination at 7q35 in the family. Thus experimental validation (in
158 patient 8401) and CNV genotyping via quantitative PCR (qPCR) was performed in all
159 individuals with DNA available from this family to assess the presence of the *CNTNAP2*
160 intronic deletion. The deletion was validated in subject 8401, and was also detected in one
161 unaffected descendant of deceased patient 8404 (Fig 1C), implying that this CNV would have
162 been present in affected subject 8404, had DNA been available. The structural variant did not
163 segregate with disease status in this family, and is unlikely to be a highly penetrant variant as
164 it was observed also in an unaffected relative (Fig 1B).

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166 *Structural variants affecting CNTNAP2 amongst psychiatric phenotypes*

167 Several deletions and duplications have been described in neuropsychiatric phenotypes thus
168 far. In Fig 2, we present a comprehensive representation of all previously reported structural
169 variants found in *CNTNAP2* in psychiatric disorders such as ASD or ID [13, 29-38],
170 schizophrenia or bipolar disorder (including the 131kb deletion found in the present study) [51-

171 54], ADHD [55], neurologic disorders such as epilepsy, Tourette syndrome or Charcot-Marie-
172 Tooth [56-60]; and finally language-related phenotypes such as speech delay, childhood
173 apraxia of speech and dyslexia [62-65]. Interestingly, the structural variants reported so far
174 frequently map in intron 1, overlapping with the 131kb deletion found in our extended family,
175 and extend in some cases up to exon 4. The distribution of those structural variants across
176 different phenotypes does not vary with those found in control populations from the database
177 of genomic variants (<http://dgv.tcag.ca/dgv/app/home>) (Fig 2), suggesting that structural
178 variants in *CNTNAP2* are not rare events associated exclusively to disease but are present with
179 rare frequency in the general population.

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181 *Analysis of CNTNAP2 common and rare variation in the susceptibility of psychiatric disorders*

182 During the last decade, several association studies have been performed to assess the role of
183 common variants of *CNTNAP2* in several psychiatric phenotypes. The functional relationship
184 between *CNTNAP2* and the language-associated transcription factor FOXP2 prompted many
185 studies to focus on language traits in autism or speech-related phenotypes [12-15, 23-28, 46-
186 48, 50], but later, additional associations were also reported in other psychiatric phenotypes
187 [41-45, 49]. In Table 1, we summarise all markers found significantly associated in these
188 previous studies, and report the corresponding *P-value* from the Psychiatric Genomics
189 Consortium GWAS for seven major psychiatric disorders: ADHD, anorexia nervosa, ASD,
190 bipolar disorder, MDD, OCD and schizophrenia. Nominal associations were found with ASD
191 for the following markers: rs802524 ($P=0.016$), rs802568 ($P=0.008$), rs17170073 ($P=0.008$),
192 and rs2710102 (which is highly correlated with 4 SNPs: rs759178, rs1922892, rs2538991,
193 rs2538976) ($P=0.036$); with schizophrenia for rs1859547 ($P=0.044$); with ADHD for
194 rs1718101 ($P=0.038$); in MDD for rs12670868 ($P=0.047$), rs17236239 ($P=0.006$), rs4431523
195 ($P=0.001$); and with anorexia nervosa for rs700273 ($P=0.013$). The nominal association at

196 rs1770073 and rs2710102 in ASD represents the only case in which the phenotype matches
197 between the original report and the PGC dataset. The two SNPs rs7794745 and rs2710102,
198 which were repeatedly reported as being associated and proposed to be functional SNPs, were
199 not strongly associated with any phenotype (the most significant signal being $P=0.036$ for
200 rs2710102 in autism). None of those associations survived corrections for multiple
201 comparisons (Table 1).

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Table 1. Common SNPs in *CNTNAP2* previously associated in psychiatric diseases, and their evidence for association in PGC datasets.

SNP	Location	Disease (Ref)	PGC Results (<i>P</i> -Value)						
			ASD	SCZ	BD	ADHD	MDD	AN	OCD
rs34712024	Promoter	ASD [25]	0.672 [^]	0.45	0.099	0.442	N/A	0.283	0.295
rs802524	Intron 1	SCZ, BD [41]	0.016 [^]	0.081	0.058	0.210	0.070	0.143	0.039
rs802568	Intron 1	SCZ, BD [41]	0.008 [^]	0.061	0.312	0.047	0.054	0.321	0.279
rs17170073	Intron 1	ASD [26]	0.008	0.903	0.558	0.883	0.306	0.031	0.101
rs1718101	Intron 1	ASD [27]	0.076 [^]	0.257	0.215	0.038	0.255	0.243	0.029
rs700273	Intron 1	ALD [42]	0.840	0.655	0.837	0.544	0.338	0.013	0.554
<u>rs7794745</u>	Intron 2	ASD [14, 23, 24]	0.906	0.734	0.498	0.393	0.173	0.877	0.503
rs10251794	Intron 3	OPN [43]	0.301	0.365	0.155	0.452	0.047 (rs12670868)	0.648	0.351
rs7804520	Intron 3	ASD [28]	0.378	0.277	0.236	0.155	0.568	0.506	0.682
rs1603450	Intron 8	LAN [15]	0.445	0.166	0.643	0.141	0.010	0.951	0.577
rs826824	Intron 9	MDD (male only) [44]	0.218	0.181	0.256	0.317	0.266	0.736	0.199
rs1859547	Intron 11	SCZ [45]	0.697	0.044	0.431	0.225	0.939	0.729	0.154
rs851715 ^{&}	Intron 13	SLI [12]	0.448	0.496	0.572	0.067	0.601	0.920	0.411
rs10246256 ^{&}	Intron 13	SLI [12, 46]	0.429	0.613	0.508	0.070	0.601 (rs851715)	0.871	0.454
<u>rs2710102</u> [#]	Intron 13	ASD, SLI, DYS, SM, ANX, LAN, MDD [12, 13, 15, 23, 46-49]	0.036	0.893	0.801	0.911	0.346	0.383	0.351
rs759178 [#]	Intron 13	SLI, LAN [12, 15]	0.037	0.890	0.799	0.929	0.332	0.363	0.347
rs1922892 [#]	Intron 13	SLI [12]	0.039	0.908	0.794	0.940	0.332 (rs759178)	0.359	0.346
rs2538991 [#]	Intron 13	SLI [12]	0.041	0.852	0.797	0.989	0.332 (rs759178)	0.366	0.338
rs17236239	Intron 13	ASD, SCZ, SLI [12, 23, 27, 46, 49]	0.142	0.290	0.278	0.883	0.006	0.622	0.954
rs2538976 [#]	Intron 13	SLI, SSD [12, 50]	0.051	0.718	0.692	0.812	0.358	0.408	0.424
rs2215798	Intron 13	ASD [26]	0.5	0.469	0.361	0.742	0.030	0.568	0.281
rs4431523	Intron 13	SLI [12]	0.275	0.844	0.676	0.614	0.001 (rs2708267)	0.933	0.972
rs2710117	Intron 14	SLI, MDD [12, 46, 49]	0.1477	0.6701	0.7566	0.2106	0.894 (rs2710121)	0.993	0.321
rs2710093	Intron 14	ASD [26]	0.4077	0.2891	0.02819	0.2943	0.090 (rs2710091)	0.8416	0.2767

223 The disease for which association at each listed SNP is given, along with the reference number
224 for each study and the approximate location of each variant within the *CNTNAP2* gene
225 structure. On the right, the *P-value* from each Psychiatric Genomics Consortium (PGC) dataset
226 is reported. Where the associated SNP was not found in the GWAS summary statistic data,
227 results for an alternative SNP are shown in parenthesis ($r^2=1$). Putative functional SNPs
228 rs7794745 and rs2710102 are underlined. No association survives correction for multiple
229 independent tests ($P < 3.8E-04$), but *P-values* < 0.05 are shown in bold. Abbreviations: ASD,
230 autism spectrum disorder; SLI, specific language impairment; DYS, dyslexia; ANX, social
231 anxiety; LAN, language in general population; SCZ, schizophrenia; BD, bipolar disorder;
232 ALD, Alcohol dependence; OPN, Openness general population; MDD, major depressive
233 disorder; SSD, speech sound disorder; N/A, SNP not genotyped; &, $r^2 > 0.97$ across the
234 following SNPs: rs851715 and rs10246256; #, $r^2 > 0.97$ across the following SNPs: rs2710102,
235 rs759178, rs1922892, rs2538991 and rs2538976; ^, summary data at this SNP was not included
236 in the latest autism GWAS (PGC2) but was present in the previous data set which included
237 5,305 ASD cases and 5,305 controls.

238 Next, we explored the contribution of common variants across *CNTNAP2* by
 239 performing a gene-based association study in European populations using GWAS summary
 240 statistics from PGC data of seven psychiatric disorders (Table 2).

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242 **Table 2. Gene-based tests for association of *CNTNAP2* across seven psychiatric disorders**
 243 **using GWAS summary statistics of the PGC data sets.**

Disease	N Cases – Controls	N SNPs tested	Gene-based P-value	Top SNP	Top SNP P- value
ADHD ^a	19,099 - 34,194	7,538	0.16	rs370840971	4.8E-04
AN ^a	3,495 - 10,982	9,318	0.33	rs138287908	8.5E-05
ASD ^a	6,197 - 7,377	5,946	0.54	rs1089600	0.0018
BD ^a	20,352 - 31,358	11,345	0.34	rs181471483	3.6E-04
MDD ^b	9,240 - 9,519	1,214	0.029	rs4725752	9.3E-04
OCD ^a	2,688 - 7,037	8,631	0.30	rs6976859	8.7E-05
SCZ ^a	33,640 - 43,456	12,264	0.11	rs78093069	1.1E-04

244 The numbers (N) of cases and controls in each dataset examined are given, along with the
 245 number of SNPs tested in each dataset. The name and *P-value* of the SNP with most significant
 246 association is given. Abbreviations: ADHD, attention-deficit/hyperactivity disorder; AN,
 247 anorexia nervosa; ASD, Autism spectrum disorder; BD, bipolar disorder; MDD, major
 248 depressive disorder; OCD, obsessive compulsive disorder; SCZ, schizophrenia; ^a, European
 249 individuals from the PGC2 data sets; ^b, European individuals from the PGC1 data sets.

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251 The test included a dense coverage of SNPs across *CNTNAP2*: from 1,214 SNPs in MDD up
 252 to 12,264 SNPs in schizophrenia. The results suggest that common variants overall do not
 253 contribute to disease susceptibility of these phenotypes (Table 2). The most significant
 254 association observed was for MDD phase 1 analysis ($P=0.029$), which is the dataset with the
 255 most modest coverage of markers.

256 In our final analysis of the PGC datasets, we selected 63 predicted functional SNPs in
 257 *CNTNAP2* and performed a cross-disorder meta-analysis, aimed to test evidence for association
 258 with common functional variants across psychiatric disorders. Nominal significance of
 259 association was observed for 11 predicted functional SNPs with *P-values* ranging from 0.01
 260 and 0.05, but none survive correction for multiple comparisons (Table 3).

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262 **Table 3. Cross psychiatric disorders meta-analysis of predicted functional SNPs.**

SNPs	Allele	Function	Datasets	I	P-val	OR
rs17480644	A/G	TFBS	ADHD, AN, BD, OCD, SCZ	8	0.083	1.039
rs1260124	A/T	TFBS	ADHD, AN, ASD, BD, OCD, SCZ	0	0.635	0.996
rs35796336	T/C	TFBS	AN, ASD, BD, OCD, SCZ	0	0.049	1.027
rs10277276	T/C	TFBS	BD, OCD, SCZ	0	0.764	0.992
rs34712024	A/G	TFBS	ADHD, AN, BD, OCD, SCZ	32	0.626	1.012
rs2462603	A/G	TFBS	ADHD, AN, ASD, BD, OCD, SCZ	0	0.303	0.993
rs1639484	A/T	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	0	0.376	1.005
rs12703814	A/G	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	0	0.965	0.999
rs1639447	A/C	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	0	0.376	0.990
rs769344	C/G	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	0	0.778	0.996
rs10280967	A/G	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	0	0.639	0.996
rs10243142	T/C	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	0	0.738	1.002
rs12535047	T/C	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ, MDD	0	0.327	0.993
rs347201	A/G	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	0	0.089	1.011
rs13234249	T/C	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ, MDD	2	0.101	1.011
rs12666908	T/C	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	0	0.914	0.999
rs11972428	T/G	SeqCons	ADHD, AN, BD, OCD, SCZ	0	0.672	1.012
rs34222835	A/G	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	4	0.045	0.979
rs10261412	A/G	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ, MDD	52	0.808	0.995
rs1826843	A/G	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ, MDD	20	0.153	0.990
rs17170356	A/G	SeqCons	ADHD, AN, BD, OCD, SCZ	0	0.137	0.972
rs4726831	A/C	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	18	0.294	0.990
rs10279700	T/C	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ, MDD	26	0.857	0.998
rs35701811	A/G	SeqCons	ADHD, AN, BD, OCD, SCZ	0	0.899	0.998
rs899617	T/C	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	0	0.026	1.014
rs747140	C/G	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	0	0.025	1.014
rs7798078	A/G	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ, MDD	0	0.091	1.011
rs34592169	A/G	Splicing	ADHD, AN, ASD, BD, OCD, SCZ	15	0.067	1.014
rs6970064	A/G	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ, MDD	7	0.462	1.004
rs17170640	A/G	SeqCons	ADHD, AN, BD, OCD, SCZ	0	0.018	1.041
rs16883690	A/C	SeqCons	BD, SCZ	66	0.673	1.066
rs7797724	T/C	SeqCons	BD, SCZ	0	0.047	1.592
rs851659	A/C	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	9	0.349	0.993
rs35815165	-/AA	SeqCons	ADHD, AN	0	0.831	1.002
rs13247212	T/C	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	0	0.797	0.996
rs1177007	A/G	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ, MDD	0	0.238	1.008
rs12154883	T/G	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	0	0.645	1.006
rs13438769	T/C	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ, MDD	0	0.057	1.018
rs2707580	T/G	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	0	0.146	0.990
rs2707581	T/C	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	0	0.156	0.990
rs2141955	A/G	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	0	0.032	1.015
rs34347668	A/C	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ	0	0.973	1.000
rs4725756	A/C	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ, MDD	0	0.015	0.984
rs2888540	T/C	SeqCons	ADHD, AN, ASD, BD, OCD, SCZ, MDD	0	0.023	1.014
rs17170789	A/T	SeqCons	ADHD, AN, BD, OCD, SCZ	0	0.566	1.010
rs17170801	A/C	SeqCons	ADHD, AN, BD, OCD, SCZ	0	0.643	0.989
rs10279343	T/C	SeqCons	ADHD, AN, BD, OCD, SCZ	1	0.568	1.014
rs1122622	A/C	SeqCons	AN, BD, OCD, SCZ	66	0.745	1.026
rs5888312	-/A	SeqCons	ADHD, AN	70	0.744	1.010
rs9648691	A/G	Splicing	ADHD, AN, ASD, BD, OCD, SCZ, MDD	64	0.806	1.002
rs987456	A/C	miRNA	ADHD, AN, ASD, BD, OCD, SCZ, MDD	29	0.570	1.004
rs2717809	C/G	miRNA	AN, BD, OCD, SCZ	0	0.746	0.989
rs2530312	A/G	miRNA	ADHD, AN, ASD, BD, OCD, SCZ	70	0.485	0.990
rs3194	A/C	miRNA	ADHD, AN, ASD, BD, OCD, SCZ, MDD	68	0.797	1.003
rs10243309	C/T	miRNA	AN, MDD	15	0.125	1.149

rs17170999	A/G	miRNA	AN, BD, OCD, SCZ, MDD	0	0.026	0.917
rs2530311	A/G	miRNA	ADHD, AN, ASD, BD, OCD, SCZ, MDD	66	0.975	0.999
rs17171000	T/C	miRNA	ADHD, AN, BD, OCD, SCZ, MDD	0	0.583	0.988
rs10251347	C/G	miRNA	ADHD, AN, BD, OCD, SCZ, MDD	0	0.428	0.986
rs2717829	C/G	miRNA	ADHD, AN, ASD, BD, OCD, SCZ, MDD	50	0.820	0.997
rs10280038	A/G	miRNA	AN, BD, OCD, SCZ, MDD	0	0.031	1.087
rs2530310	T/C	miRNA	ADHD, AN, ASD, BD, OCD, SCZ, MDD	67	0.666	0.994
rs17171006	T/C	miRNA	ADHD, AN, BD, OCD, SCZ	0	0.493	0.985

263 For each predicted functional SNP, the alternative alleles and predicted function are listed. *P*-
 264 *values* were calculated considering fixed-model effect, except SNPs with evidence of
 265 heterogeneity ($I > 50$) where odds ratios (OR) were considered under random-effects.
 266 Nominally significant associations are indicated in bold (P -*values* < 0.05), but none exceed
 267 correction for multiple testing ($P < 7.9E-04$). Abbreviations: TFBS, transcription factor binding
 268 site; SeqCons, sequence conserved nucleotide across species; miRNA, predicted miRNA
 269 binding site; Splicing, exonic splicing enhancer (ESE).
 270

271 The only SNP predicted to be functional and which was previously reported as being associated
 272 with autism was rs34712024 (Table 2) [25], but this variant was not associated with autism in
 273 PGC dataset ($P=0.67$), nor other psychiatric phenotypes examined (Table 2).

274 *De novo* variants in protein-coding genes which are predicted to be functionally
 275 damaging are considered to be highly pathogenic and have been extensively explored to
 276 implicate genes in psychiatric diseases, especially in ASD and schizophrenia [67]. We explored
 277 publically available sequence data from previous projects in psychiatric disorders to assess the
 278 rate of coding *de novo* variants in *CNTNAP2* using two databases (NPdenovo,
 279 <http://www.wzgenomics.cn/NPdenovo/>; and denovo-db, [http://denovo-](http://denovo-db.gs.washington.edu/denovo-db/)
 280 [db.gs.washington.edu/denovo-db/](http://denovo-db.gs.washington.edu/denovo-db/)). No truncating or missense variants were identified across
 281 *CNTNAP2* in 15,539 families (including 2,163 controls), and synonymous variants were
 282 reported in only two probands with developmental disorder (Table 4).

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288 **Table 4. *CNTNAP2* de novo variants identified across several disease-specific sequencing**
289 **projects.**

Phenotype	N Families	Intronic	Synonymous	Missense
ASD	6,171	106	-	-
SCZ	1,164	-	-	-
EE	647	-	-	-
ID	1,101	-	-	-
DD	4,293	-	2	-
Controls	2,163	13	-	-

290 The number (N) of families in each dataset examined is given. The full list of *de novo* variants
291 observed is listed in S2 Table. Abbreviations: ASD, autism spectrum disorder; SCZ,
292 schizophrenia; EE, epilepsy; ID, intellectual disability; DD, developmental disability.

293

294 Finally, we explored the potential impact of pathogenic ultra-rare variants (URV) in
295 *CNTNAP2* using available sequencing datasets of 4,483 patients with ASD and 6,135 patients
296 with schizophrenia compared with 13,042 controls. We considered only those variants
297 predicted to be pathogenic in both SIFT and Polyphen and which are ultra-rare (MAF<0.0001
298 in Non-Finnish European population). No difference in the total number of URV was observed
299 between ASD and controls ($P=0.11$), or between schizophrenia patients and controls ($P=0.78$)
300 (Table 5).

301

302 **Table 5. Burden analysis of *CNTNAP2* ultra-rare variants (URVs) in ASD and SCZ.**

	N Individuals	N Pathogenic URVs	<i>P-Value</i>
Controls	13,042	59	
SCZ	6,135	26	0.78
ASD	4,483	29	0.11

303 The selection of variants included missense variants which are predicted to be pathogenic,
304 truncating variants and canonical splice-site variants. The full list of URVs observed is
305 provided in S3 Table. Abbreviations: SCZ, schizophrenia; ASD, autism spectrum disorder

306 **DISCUSSION**

307 During the last decade, the *CNTNAP2* gene has received considerable attention in the
308 psychiatric genetics field, with a large body of studies examining gene dosage, common or rare
309 variants across multiple major psychiatric disorders, which together provided compelling
310 evidence that *CNTNAP2* may be a risk gene with pleiotropic effects in psychiatry. While
311 homozygous mutations in this gene lead to a rare and severe condition described as CASPR2
312 deficiency disorder (CDD) [7], characterized by profound intellectual disability, epilepsy,
313 language impairment or regression [7, 8], heterozygous mutations or common variants have
314 been suggested to be implicated in autism, whose features overlap with some observed in CDD.
315 *CNTNAP2* is categorised in the SFARI database (<https://gene.sfari.org>) as a strong candidate
316 gene for ASD (category 2.1). Heterozygous deletions encompassing the *CNTNAP2* gene were
317 described not only in autism but also in a wide range of phenotypes, including psychiatric or
318 neurologic disorders, and language-related deficiencies. These structural variants were
319 generally described as causative or highly penetrant [13, 29, 31, 55, 57, 59].

320 Here, we describe a new deletion in a bipolar disorder patient encompassing intron 1 of
321 *CNTNAP2*, which overlaps with structural variants described in a number of other psychiatric
322 patients. This heterozygous deletion, which removes the FOXP2 transcription factor binding
323 site, was found in an individual with bipolar I disorder from an extended family with five
324 affected members. This deletion was observed in only two of the affected relatives and was
325 absent from two affected relatives, but was also observed in one unaffected relative who
326 underwent diagnostic interview at age >40 and therefore beyond the typical age of symptom
327 onset. Hence, the deletion was not segregating with the disease and is unlikely to represent a
328 highly penetrant risk variant in this family. Examination of the distribution of all structural
329 variants described thus far in psychiatric or neurologic patients showed comparable mapping
330 with those found in the general population, suggesting that structural variants affecting

331 *CNTNAP2* may be less relevant in disease susceptibility than previously considered. Eleven
332 CNVs are described in the general population against sixteen expected ($z=0.43$) in ExAC
333 database (<http://exac.broadinstitute.org>), and the haploinsufficiency score (0.59) is relatively
334 moderate [68], suggesting that *CNTNAP2* has a moderate tendency to be intolerant to structural
335 variants. However, a case-control CNV analysis is needed in psychiatric disorders, but would
336 require a very large sample due to the rarity of CNVs at this locus. A close clinical psychiatric
337 examination of the 66 parents with heterozygous deletions across *CNTNAP2* of CDD provide
338 information on the prevalence of psychiatric conditions in individuals carrying *CNTNAP2*
339 CNVs. All heterozygous family members carrying deletions or truncating mutations were
340 described as phenotypically healthy, suggesting a lack of correlation between these deletions
341 and any major psychiatric condition. Furthermore, parents who were carriers for heterozygous
342 deletions in psychiatric/neurologic patients were described as unaffected at the time of
343 reporting [13, 29, 31, 37, 54, 62], with the exception of one father of a proband with neonatal
344 convulsion or another father of an epileptic patient reported as affected [56, 59]. Moreover,
345 discordant segregation for deletions in *CNTNAP2* was also observed in an ASD sib-pair [13].
346 Several psychiatric patients who were reported to carry heterozygous structural variants in
347 *CNTNAP2* were also described with translocations or other chromosomal abnormalities [29,
348 30, 33, 34, 56, 58, 62-65], therefore it is possible that these aberrations may explain the
349 phenotype independently from the observed CNVs in *CNTNAP2*.

350 *CNTNAP2*^{-/-} knock-out mice have been proposed as valid animal model for ASD
351 considering the phenotypic similarities between ASD and the CASPR2 deficiency disorder [2].
352 *CNTNAP2*^{-/-} knock-out mice showed abnormalities in the arborisation of dendrites, maturation
353 of dendritic spines, defects in migration of cortical projection neurons, and reduction of
354 GABAergic interneurons [2, 4]. Controversially, ASD is not a core feature in the most recent
355 patient series reported with CASPR2 deficiency disorder [7, 8]. The association previously

356 proposed around the relationship between heterozygous deletions in *CNTNAP2* and ASD does
357 not have a support from mouse models, as heterozygous mice did not show any behavioural or
358 neuropathological abnormalities that were observed in homozygous knockouts [2].
359 Notwithstanding this, it is possible that the combination of heterozygous *CNTNAP2* deletions
360 in a genomic background of increased risk (through inheritance of other common and rare risk
361 variants at other loci) may lead to psychiatric, behavioural or neuropathological abnormalities.

362 Common variants in *CNTNAP2* are another class of genetic variation associated with
363 several psychiatric or language-related phenotypes. The most interesting finding from these
364 studies converge on markers rs7794745 and rs2710102, originally reported in ASD [13, 14],
365 and replicated later in ASD or implicated in other phenotypes [12, 15, 23, 24, 46-48].
366 Neuroimaging studies have supported the notion that these common variants play a role in
367 psychiatric disorders. SNP rs2710102 has been implicated in brain connectivity in healthy
368 individuals [16, 18, 19], and rs7794745 was implicated in audio-visual speech perception [69],
369 voice-specific brain function [22], and was associated with reduced grey matter volume in left
370 superior occipital gyrus [20, 21]. These studies focused principally on language tasks in general
371 population, given the reported suggestive implications of *CNTNAP2* in language impairment
372 traits of ASD or language-related phenotypes. However, the direct role of *CNTNAP2* in
373 language is still unclear; indeed the language regression observed in patients with CASPR2
374 deficiency are concomitant with seizure onset and may represent a secondary phenotypic effect
375 caused by seizures [7]. On the other hand, the first genetic association of rs7794745 and
376 rs2710102 with ASD, as well as the other psychiatric diseases were based in studies with
377 limited sample size, and recent studies failed to replicate associations between the two markers
378 and ASD [70, 71]. Individual alleles associated in the past with limited numbers of patients
379 warrant replications in adequately powered samples to ascertain *bona fide* findings considering
380 the small size effects of common variants [72], which we attempted here using the largest case-

381 control cohorts currently publicly available (PGC datasets). We did not find evidence for
382 significant association of previous reported common variants, nor did we find functional SNPs
383 with a role across disorders, or observe a combined effect for common variants of *CNTNAP2*
384 in the susceptibility of psychiatric disorders.

385 Rare variants of *CNTNAP2* both in the promoter or coding region were also reported to
386 play a role in the pathophysiology of ASD [25, 33]. A recent study including a large number
387 of cases and controls did not find association of rare variants of *CNTNAP2* in ASD [73]. Here
388 we report the largest sample investigated thus far in ASD and schizophrenia for rare variants
389 in *CNTNAP2*, which suggest that rare variants from this gene do not play a major role in these
390 two psychiatric disorders. Furthermore, the identification of *de novo* variants in *CNTNAP2* in
391 combined psychiatric sequencing projects of over 15,500 trios suggest that *de novo* variants in
392 this gene do not increase risk for psychiatric disorders.

393 While functional studies show a relationship between certain deletions or rare variants
394 of *CNTNAP2* with neuronal phenotypes relevant to psychiatric illness [25, 54, 74], we show
395 that the genetic link between these variants and psychiatric phenotypes is tenuous. However,
396 this does not dispel the evidence that the *CNTNAP2* gene, or specific genetic variations within
397 this gene, may have a real impact on neuronal functions or brain connectivity.

398 Nowadays we are able to combine large datasets to ascertain the real impact of
399 candidate genes described in the past in psychiatric disorders. Here we performed analyses
400 using large publically available datasets investigating a range of mutational mechanisms which
401 impact variability of *CNTNAP2* across several psychiatric disorders. In conclusion, our results
402 converge to show a limited or likely neutral role of *CNTNAP2* in the susceptibility of
403 psychiatric disorders.

404

405

406 MATERIALS AND METHODS

407 *Extended family with bipolar disorder and CNV in CNTNAP2*

408 The extended family presented here (Fig 1B) provides a molecular follow-up from a previously
409 reported whole exome sequencing (WES) study of multiplex BD families, augmented with
410 CNV microarray data [66]. This multigenerational pedigree, was collected through the Mood
411 Disorders Unit and Black Dog Institute at the Prince of Wales Hospital, Sydney, and the School
412 of Psychiatry (University of New South Wales in Sydney) [75-79]. Consenting family
413 members were assessed using the Family Interview for Genetic Studies (FIGS) [80], and the
414 Diagnostic Interview for Genetic Studies (DIGS) [81]. The study was approved by the Human
415 Research Ethics Committee of the University of New South Wales, and written informed
416 consent was obtained from all participating individuals. Blood samples were collected for DNA
417 extraction by standard laboratory methods. Three of the five relatives with bipolar disorder
418 type I (BD-I) had DNA and WES-derived genotype data available, and six unaffected relatives
419 with DNA and WES data were available for haplotype phasing and segregation analysis (Fig
420 1B).

421 Genome-wide CNV analysis was performed via CytoScan® HD Array (Affymetrix,
422 Santa Clara, CA, USA) in 2 distal affected relatives (individuals 8410 and 8401; Fig 1B), using
423 the Affymetrix Chromosome Analysis Suite (ChAS) software (ThermoFisher, Waltham, MA,
424 USA). Detailed information on CNV detection and filtering criteria have been previously
425 described [66]. We identified a 131kb deletion in intron 1 of *CNTNAP2* in individual 8401.
426 WES-derived genotypes were used for haplotype assessment to infer CNV segregation
427 amongst relatives, as previously described [66]. Next, we experimentally validated the
428 *CNTNAP2* CNV via quantitative PCR (qPCR) in all available family members. Validation was
429 performed in quadruplicate via a SYBR Green-based quantitative PCR (qPCR) method using
430 two independent amplicon probes, each compared with two different reference amplicon

431 probes in the *FOXP2* and *RNF20* genes (S1 Table). Experimental details are available upon
432 request.

433

434 *Common variant association in CNTNAP2 using publically available datasets*

435 We sought to replicate previously reported *CNTNAP2* SNP associations in a range of
436 psychiatric phenotypes or traits using GWAS summary-statistic data of the Psychiatric
437 Genomics Consortium ([https:// med.unc.edu/pgc/results-and-downloads](https://med.unc.edu/pgc/results-and-downloads)).

438 Firstly, we report the corresponding *P-values* of specific previously associated markers
439 for case-control cohorts with autism spectrum disorder (ASD), schizophrenia (SCZ), bipolar
440 disorder (BD), attention-deficit hyperactivity-disorder (ADHD), major depressive disorder
441 (MDD), anorexia nervosa (AN), and obsessive compulsive disorder (OCD). If a specific SNP
442 marker was not reported in an individual GWAS dataset, we selected another marker in high
443 linkage disequilibrium ($r^2 \sim 1$, using genotype data from the CEU, TSI, GBR and IBS European
444 populations in 1000genomes project; <http://www.internationalgenome.org>).

445 Next, a gene-based association for common variants was calculated with MAGMA
446 [82], using variants within a 5 kb window upstream and downstream of *CNTNAP2*. Selected
447 datasets were of European descent, derived from GWAS summary statistics of the Psychiatric
448 Genomics Consortium (<https://med.unc.edu/pgc/results-and-downloads>): SCZ (33,640 cases
449 and 43,456 controls), BD (20,352 cases and 31,358 controls), ASD (6,197 and 7,377 controls),
450 ADHD (19,099 cases and 34,194 controls) and MDD (9,240 cases and 9,519 controls) [83-87].
451 Analyses were performed combining two different models for higher statistical power and
452 sensitivity when the genetic architecture is unknown: the combined *P-value* model, which is
453 more sensitive when only a small proportion of key SNPs in a gene show association; and the
454 mean SNP association, which is more sensitive when allelic heterogeneity is greater, and a
455 larger number of SNPs show nominal association.

456 Finally, we selected SNPs predicted to be functional within a 5kb window
457 upstream/downstream of *CNTNAP2* (e.g. located in transcription factor binding sites, miRNA
458 binding sites etc; <https://snpinfo.niehs.nih.gov>), and assessed a potential cross-disorder effect
459 using GWAS summary statistics data of the PGC by performing a meta-analysis in PLINK
460 [88]. The Cochran's Q-statistic and I^2 statistic were calculated to examine heterogeneity among
461 studies. The null hypothesis was that all studies were measuring the same true effect, which
462 would be rejected if heterogeneity exists across studies. For all functional SNPs, when
463 heterogeneity between studies was $I > 50\%$ ($P < 0.05$), the pooled OR was estimated using a
464 random-effects model.

465

466 *Analysis of rare variants in CNTNAP2 in ASD and schizophrenia, and de novo variants across*
467 *psychiatric cohorts*

468 The impact of rare variants of *CNTNAP2* was assessed using sequencing-level data from the
469 following datasets: WES from the Sweden-Schizophrenia population-based Case-Control
470 cohort (6,135 cases and 6,245 controls; dbGAP accession: phs000473.v2.p2); ARRA Autism
471 Sequencing Collaboration (490 BCM cases, BCM 486 controls, and 1,288 unrelated ASD
472 probands from consent code c1; dbGAP accession: phs000298.v3.p2); Medical Genome
473 Reference Bank (2,845 healthy Australian adults; <https://sgc.garvan.org.au/initiatives/mgrb>);
474 individuals from a Caucasian Spanish population (719 controls [89, 90]); in-house ASD
475 patients (30 cases; [91]); and previous published data set in ASD (2,704 cases and 2,747
476 controls [73]). The selection of potentially etiologic variants was performed based on their
477 predicted pathogenicity (missense damaging in both SIFT and polyphen 2, canonical splice
478 variants, stop mutation and indels) and minor allele frequency (MAF < 0.0001 in non-Finnish
479 European populations using the Genome Aggregation Database;
480 <http://gnomad.broadinstitute.org/>). A chi square statistic was used to compare separately the

481 sample of schizophrenia patients (6,135 cases) and the combined ASD data sets (4,512 cases)
482 with the combined control data sets (13,042 individuals).

483 Two databases for *de novo* variants were used to identify *de novo* variants in *CNTNAP2*
484 [92, 93], which comprise data for the following samples: autism spectrum disorder (6,171
485 families), schizophrenia (1,164 families), epilepsy (647 families), intellectual disability (1,101
486 families), developmental disorders (4,293 families) and controls (2,163).

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501 **ACKNOWLEDGMENTS**

502 This study was funded by the Australian National Medical and Health Research Council
503 (NHMRC) Project Grant 1063960 and supported by NHMRC Project Grant 1066177, and
504 Program Grant 1037196. We gratefully acknowledge the Janette Mary O’Neil Research
505 Fellowship (to JMF) and Mrs. Betty Lynch for supporting this work. DNA for the bipolar
506 disorder family was extracted by Genetic Repositories Australia (GRA;
507 www.neura.edu.au/GRA), an Enabling Facility that was supported by NHMRC Enabling Grant
508 401184. DNA samples were prepared and whole exome sequenced at the Lottery State
509 Biomedical Genomics Facility, University of Western Australia. We would also thank Xose S.
510 Puente from the University of Oviedo (Spain) for providing us the data from a Spanish control
511 population. We are grateful to all participants and their families, as well as clinical
512 collaborators who were originally involved in collecting and phenotyping these families,
513 including Laila Tabassum, and Adam Wright (UNSW).

514

515 **FINANCIAL DISCLOSURES**

516 The authors report no biomedical financial interests or potential conflicts of interest.

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852

853 **FIGURES**

854

855 **Fig 1. CNV deletion encompassing intron 1 of *CNTNAP2* in an extended family with**
856 **bipolar disorder.** A) CytoScan HD array output image shows the position of the drop in signal
857 intensity of 340 probes, indicating a deletion spanning 131kb (chr7:146203548-146334635;
858 GRCh37/hg19) found in the patient 8401. The position of the FOXP2 binding site within the
859 deletion is shown above. B) The bipolar pedigree includes five patients with bipolar disorder I
860 (BPI) across two generations. Symbols: _, individuals with DNA available; &, individuals with
861 whole exome data; #, individuals analysed for genome-wide CNVs through the CytoScan HD
862 array; blue squares, individuals included in CNV qPCR validation and genotyping analysis, for
863 which heterozygous deletion carriers are indicated as “+/del” and non-carriers are indicated as
864 “+/+”. Inferred genotypes are in parentheses. C) Gene dosage results of the qPCR experiments
865 validating the deletion in patient 8401, and showing the deletion in unaffected subject 8407.

866 **Fig 2. Overview of heterozygous CNVs spanning the *CNTNAP2* gene across several**
867 **diseases.** Abbreviations: ID (Intellectual disability), ASD (autism spectrum disorder), SCZ
868 (schizophrenia), BD (bipolar disorder), ADHD (Attention-deficit/hyperactivity disorder), EP
869 (epilepsy), TS (Tourette syndrome), CMT2 (axonal Charcot-Marie-Tooth), and SS (Speech
870 spectrum: speech delay, childhood apraxia of speech and dyslexia). In parenthesis is reported
871 the reference to each study. PS refers to this present study. *, additional rearrangements
872 reported in this patient. The dashed lines represent the exons and the upper box shows the
873 position of the FOXP2 binding site. In dark shading, CNVs \geq 80kb found in the general
874 populations from the Database of Genomic Variants are shown.

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886 **SUPPORTING INFORMATION**

887 **S1 Table. Primers used in the CNV validation for the *CNTNAP2* intronic deletion.**

888 **S2 Table. Full list of *de novo* variants in *CNTNAP2* gene.**

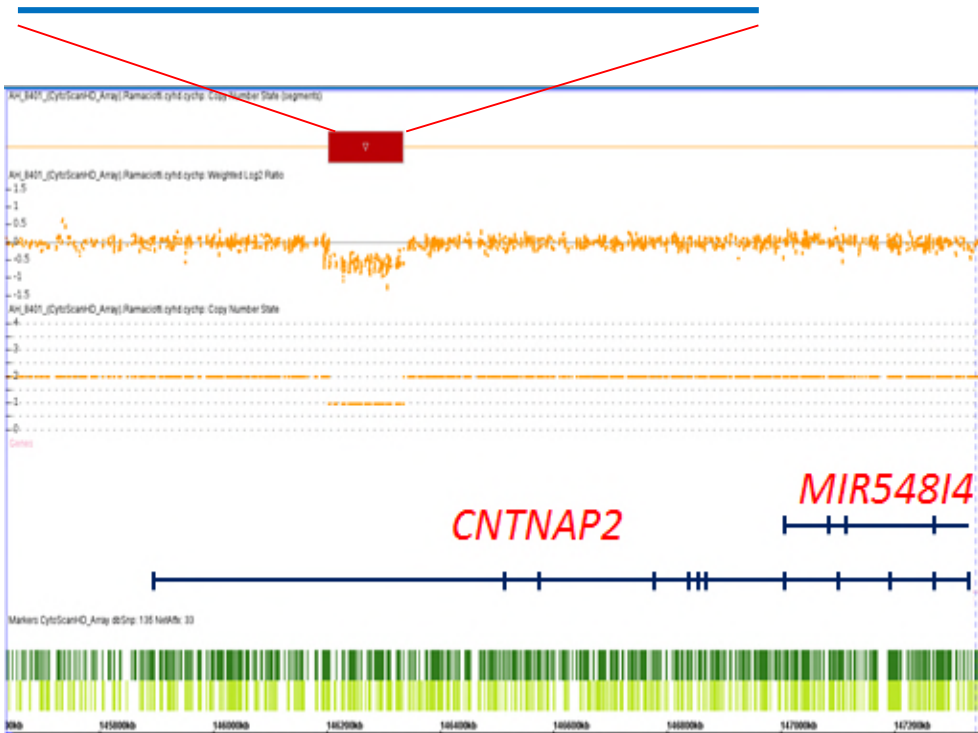
889 **S3 Table. Full list of Ultra-Rare Variants (URVs) in available sequencing datasets.**

890

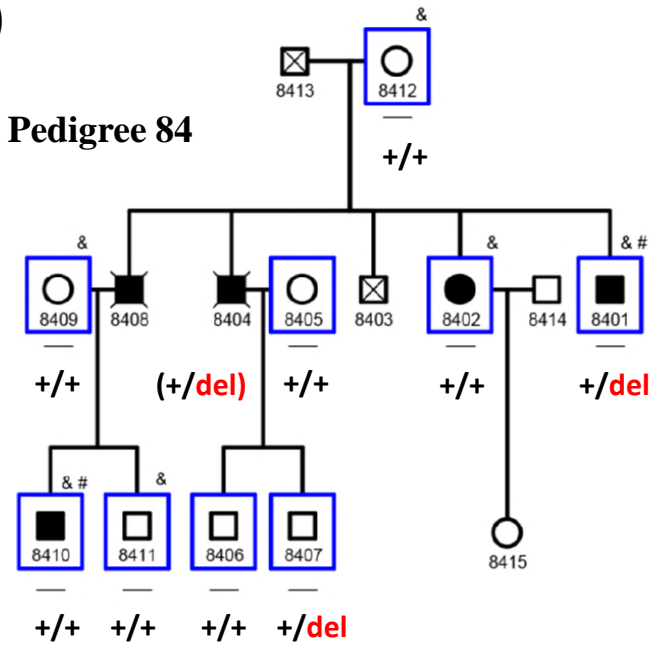
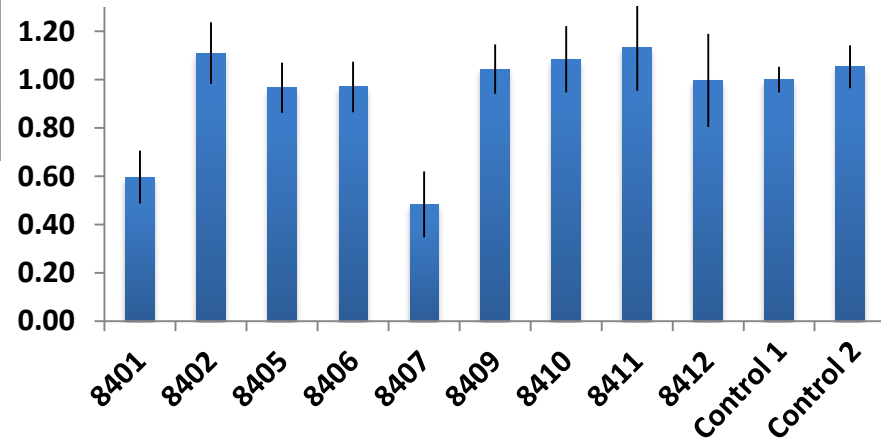
A)

FOXP2-binding site (chr7:146215016-146215040)
 AGCTGCTTTCAAATTTAAGCAATCAAGTG

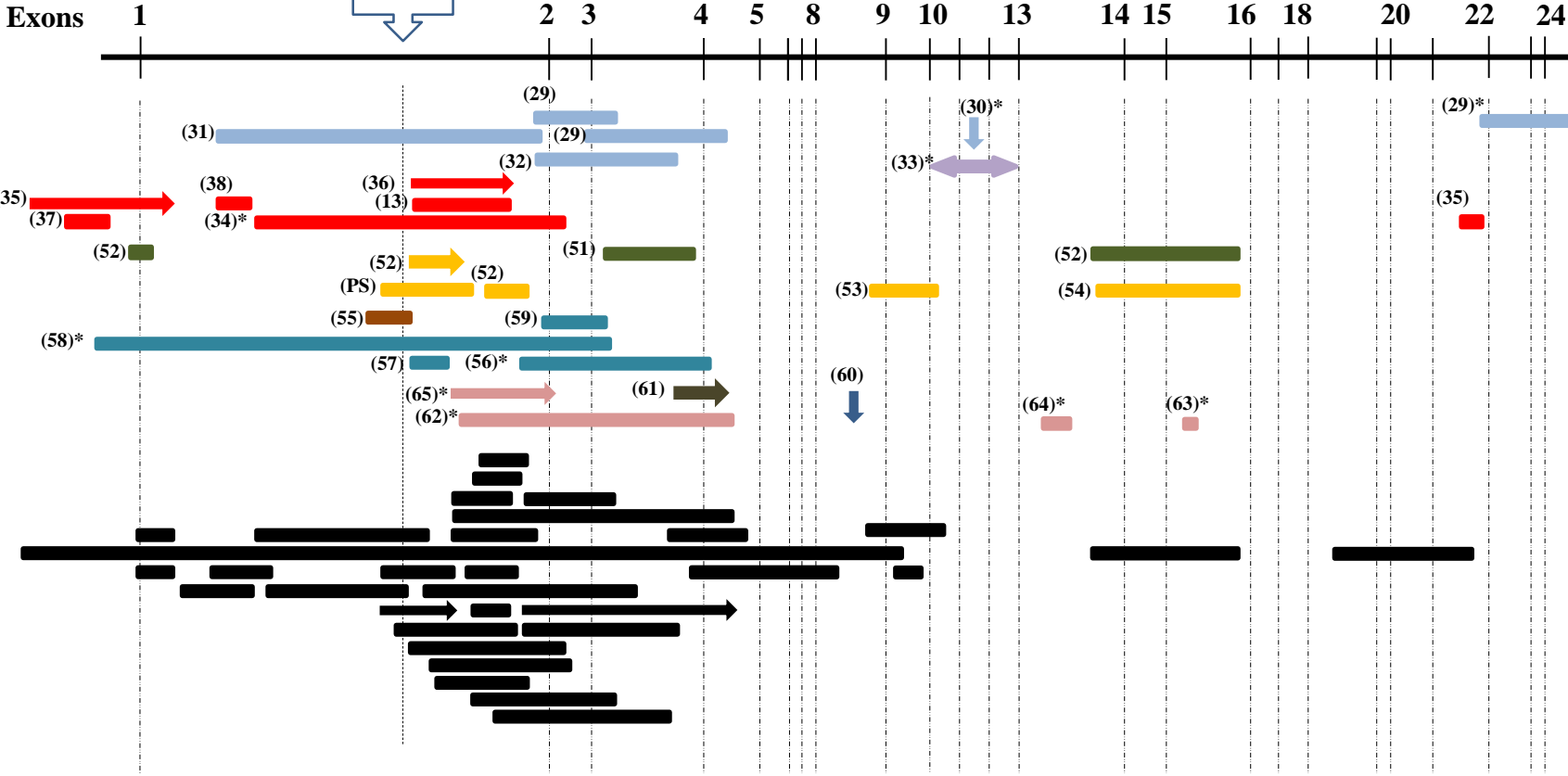
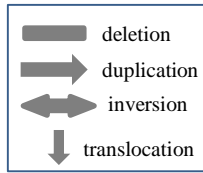
131kb (340 probes)

**B)**

Pedigree 84

**C)**

CNTNAP2



- ID (29-32)
- ⇔ ASD-ID (33)
- ⇨ ASD (13,34-38)
- SCZ (51,52)
- ⇨ BD (PS,52-54)
- ADHD (55)
- ⇨ EP (56-59)
- ⇩ TS (60)
- CMT2 (61)
- ⇨ SS (62-65)

- DGV (CNV>80kb)