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Rare mutations in N-methyl-D-aspartate glutamate receptors in autism spectrum disorders and schizophrenia

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Pharmacological, genetic and expression studies implicate N-methyl-D-aspartate (NMDA) receptor hypofunction in schizophrenia (SCZ). Similarly, several lines of evidence suggest that autism spectrum disorders (ASD) could be due to an imbalance between excitatory and inhibitory neurotransmission. As part of a project aimed at exploring rare and/or *de novo* mutations in neurodevelopmental disorders, we have sequenced the seven genes encoding for NMDA receptor subunits (NMDARs) in a large cohort of individuals affected with SCZ or ASD ($n = 429$ and 428 , respectively), parents of these subjects and controls ($n = 568$). Here, we identified two *de novo* mutations in patients with sporadic SCZ in *GRIN2A* and one *de novo* mutation in *GRIN2B* in a patient with ASD. Truncating mutations in *GRIN2C*, *GRIN3A* and *GRIN3B* were identified in both subjects and controls, but no truncating mutations were found in the *GRIN1*, *GRIN2A*, *GRIN2B* and *GRIN2D* genes, both in patients and controls, suggesting that these subunits are critical for neurodevelopment. The present results support the hypothesis that rare *de novo* mutations in *GRIN2A* or *GRIN2B* can be associated with cases of sporadic SCZ or ASD, just as it has recently been described for the related neurodevelopmental disease intellectual disability. The influence of genetic variants appears different, depending on NMDAR subunits. Functional compensation could occur to counteract the loss of one allele in *GRIN2C* and *GRIN3* family genes, whereas *GRIN1*, *GRIN2A*, *GRIN2B* and *GRIN2D* appear instrumental to normal brain development and function.

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Introduction

Glutamate is the major excitatory neurotransmitter of the central nervous system and is implicated in many basic neuronal functions and central nervous system processes, in particular, learning, memory and synaptic plasticity. N-methyl-D-aspartate receptors (NMDAR) are voltage-dependant ionotropic glutamate receptors and are tetramers containing two NR1 subunits in association with two NR2 (A, B, C or D) and/or NR3 (A or B) subunits. Many of the physiological and pharmacological properties of the NMDAR depend on the specific NR2 and NR3 subunits composition.¹ Behavioral tasks that involve NMDAR include associative learning, working memory, behavioral flexibility or attention. Thus, NMDAR signaling could be a point of convergence to explain

characteristic patterns of symptoms and neurocognitive deficits observed in neurodevelopmental disorders.

In schizophrenia (SCZ), dysregulation of brain glutamate transmission through NMDAR has been proposed as an etiological factor. Indeed, NMDAR antagonists (e.g., ketamine, phencyclidine) produce behavioral and cognitive deficits in normal subjects that closely mimic SCZ,^{2–4} and patients with anti-NR1 encephalitis display many SCZ-like symptoms and/or loss of memory.^{5–7} Mutant mice lacking NR1 subunit in about half of cortical interneurons early in post-natal development exhibit anxiety-like behaviors and working memory deficits, emerging at adolescence.⁸ In addition, NMDAR density and NMDAR subunit composition are reported abnormal in post-mortem brains of patients with SCZ.^{9–15}

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SCZ has been associated with genetic variants in some of the NMDAR subunit (*NMDARs*) genes including *GRIN2A*, *GRIN2B* and *GRIN1*, although contradictory results were reported (see the SCZgene database: <http://www.szgene.org/>).¹⁶ In addition, several genes that have been associated with increased risk for SCZ, including *NRG1-erb4*, *serine racemase*, *DAO*, *G72* or *dysbindin-1*, are implicated in the modulation of NMDAR activity.^{17–19} Lastly, the only consistent report from independent genome-wide association studies in SCZ is the association with genetic markers in the major histocompatibility complex region,^{20–22} and recent evidence suggests that major histocompatibility complex region is implicated in neuronal plasticity²³ and glutamatergic receptor modulation.²⁴

Dysfunction of NMDA signaling has also been implicated in a wide range of neurological or neurodevelopmental disorders such as intellectual disability (ID),^{25,26} in which it appears that specific NMDARs may have different roles in synaptic plasticity and excitotoxicity.²⁷ In autism spectrum disorders (ASD), potentially causative mutations in *NMDARs* genes have been recently reported,^{28,29} and a number of ASD animal models were associated with NMDA abnormalities.^{30–33} Moreover, disruption in the balance between excitation and inhibition is a commonly proposed disease mechanism for ASD,³⁴ and an open-label study with memantine, an antagonist of the NMDAR, significantly improved symptoms.³⁵ Altogether, the literature suggests that NMDAR number and composition have a crucial role in neurodevelopment and cognition.

Recent studies indicate that rare and penetrant *de novo* mutations could account for some cases of SCZ. For examples, rare chromosomal abnormalities are more frequent in SCZ than in controls, especially rare *de novo* copy number variations^{22,36,37} similar to what had been reported in other neurodevelopmental disorders such as ASD³⁸ and ID.^{8,39} In addition, recent reports, including ours, issued from the Synapse to Disease Project, support the implication of *de novo* single nucleotide mutations in ID,^{25,40–42} ASD^{29,41,43} or SCZ.^{44–47} Interestingly, among rare *de novo* variants reported associated with SCZ, two affect genes that may be linked to the glutamatergic synapses: *SHANK3*, encoding a scaffolding protein at the excitatory synapse and *KIF17* encoding a synaptic motor protein, involved in the transport of NMDA receptor 2B (NR2B) to the synapse.⁴⁸

As part of the Synapse to Disease Project, in which we screened 600 synaptic genes, we screened all exons of the seven NMDARs in 571 individuals, including 429 patients with SCZ and 142 patients with ASD. We also screened all exons of the NMDARs (except for *GRIN3B* and *GRIN1*) in 283 healthy controls. Further, we screened additional cohort of ASD patients ($n=286$) for *GRIN2B*, *GRIN2C* and *GRIN3B*, and additional cohort of controls from the general population ($n=285$) for *GRIN2C* and *GRIN3A*. For *GRIN1*, we screened only this general population cohort of controls. Finally, we determined transmission of all unique variants, identified in only one individual, by sequencing parents of the carrier (292 SCZ and 404 ASD patients, with 2 parents available for screening).

We here report the identification of *de novo* mutations in patients with SCZ in the *GRIN2A* gene and in a patient with ASD in the *GRIN2B* gene. Also, we report the absence of truncating mutation in the *GRIN1*, *GRIN2A*, *GRIN2B* and

GRIN2D genes in patients or controls, whereas we identified truncating mutations in the *GRIN2C*, *GRIN3A* and *GRIN3B* genes in patients and controls.

Methods

Subject collection

SCZ cohort. Subjects with SCZ were selected from over 1000 families ascertained for genetic studies, for which DNA samples were available. The screened cohort included 429 subjects with SCZ. We selected in priority patients with no family history of psychiatric disorders ($n=188$ patient) and patients for which both parents were available ($n=292$ patients with both parents). To ensure accurate diagnoses, all individuals were evaluated by experienced investigators (JR, LD, MOK and RJ) using the Diagnostic Interview for Genetic Studies or the Kiddie Schedule for Affective Disorders and SCZ and multidimensional neurological, psychological, psychiatric and pharmacological assessments in the recruiting centers. Family history for psychiatric disorders was also collected using the Family Interview for Genetic Studies. All Diagnostic Interview for Genetic Studies and Family Interview for Genetic Studies have been reviewed by two or more psychiatrists for a final consensus diagnosis, based on Diagnostic and Statistical Manual of Mental Disorders (DSM)-III-R or DSM-IV at each center. For all probands, specific inclusion criteria for the present study were as follows: (1) the selected proband was definitely affected with SCZ only, not schizophreniform psychosis or bipolar disorders with psychosis; (2) in families with multiple affected individuals, we selected the most severe SCZ case with early (<18 years) or childhood onset (<12 years), and/or additional neurodevelopmental problems, such as ID, dyslexia and epilepsy, but not autistic disorder; for the childhood onset SCZ cohort, ID and epilepsy were exclusionary; (3) Family history was well documented. Finally, exclusion criteria included patients with psychotic symptoms mainly caused by alcohol, drug abuse, or other clinical diagnoses including cytogenetic abnormalities. The final panel of the samples of 429 SCZ subjects were recruited from different centers, that is, 1) 28 cases with childhood-onset SCZ and 66 with later onset SCZ from Judith L Rapoport (NIMH, Bethesda, MD, USA). Cases with childhood onset SCZ were recruited in USA nationwide and assessed as previously described.⁴⁹ Individuals in this cohort known to carry the Velo-Cardio-Facial Syndrome (VCFS) deletion on chromosome 22q11 were excluded; 2) 38 cases from Marie-Odile Krebs (Paris, France);⁵⁰ 3) 84 cases from Lynn E De Lisi;⁵¹ 4) 7 cases of adult onset SCZ selected from each of seven large highly consanguineous pedigrees with ≥ 10 affected individuals with SCZ and schizoaffective disorders from Pakistan; 5) 144 cases from Ridha Joober (Montreal, QC, Canada) including 39 cases from Tunisia;⁵² and 6) 62 cases from Hungary.⁵³ The detailed description of recruitment and ascertainment strategy, diagnostic instruments and criteria have been reported by each center in their previous publications.

ASD cohort. The final panel of samples included 428 autistic patients (404 with parental DNA of both parents available).

Diagnostic and selection criteria for the ASD subjects are described in detail elsewhere.⁵⁴ We screened 142 autistic patients for the seven *NMDAR* genes, and 286 additional cases were screened for variants in *GRIN2B*, *GRIN2C* and *GRIN3B*. Briefly, all subjects were diagnosed using DSM-IV or DSM-III-R criteria, and depending of the recruitment site, Autism Diagnostic Interview-Revised and the Autism Diagnostic Observation Schedule were used. In addition, the Autism Screening Questionnaire was also completed for all subjects. We excluded patients with an estimated mental age < 18 months, a diagnosis of Rett syndrome or childhood disintegrative disorder, and patients with evidence of any psychiatric and neurological conditions including birth anoxia, rubella during pregnancy, fragile-X disorder, encephalitis, phenylketonuria, tuberous sclerosis, Tourette and West syndromes.

Control cohorts. Blood samples were obtained from a total of 568 control subjects. The first control series comprised healthy volunteers that were screened and selected using standardized procedures. Only individuals without any neuropsychiatric symptoms or family history of neuropsychiatric problems were included as SCZ-negative controls. These included 283 adult subjects (131 males; 152 females), 225 of whom were of European origin and 58 were non-Europeans (Tunisian). The second group of controls comprised 285 subjects from a general population control group of mainly French Canadian origin, for whom we also had parental DNA (Quebec Newborn Twin Study; 150 males, 135 females).⁵⁵ We have systematically sequenced all exons of the seven *NMDAR* genes (except for *GRIN3B* and *GRIN1*) in the cohort of healthy volunteer controls. When rare truncating mutations were identified, we further screened all exons in additional controls from general population (Quebec Newborn Twin Study). For *GRIN1*, we screened only this general population cohort of controls.

DNA preparation. Genomic DNA was extracted from blood using Puregene extraction kit (Gentra System, Minneapolis, MN, USA). For certain individuals in whom blood DNA was limiting, we used DNA isolated from a lymphoblastoid cell line derived from the individual for the screen. In all cases, rare mutations were confirmed using blood-derived DNA to rule out variations having arisen during production or growth of the lymphoblastoid cell line.

Paternity testing. Paternity, maternity and unique genetic identification of each individual of all families (subject families and controls) was confirmed using 5–14 highly informative unlinked microsatellite markers as described in Gauthier *et al*.⁴⁶

Gene screening, variation analysis and bioinformatics. In each proband, we sequenced the coding region and the splice junction of *GRIN1* (chr9: 140,033,609–140,063,207, hg19 assembly), *GRIN2A* (chr16: 9,847,267–10,276,263), *GRIN2B* (chr12: 13,714,410–14,133,022), *GRIN2C* (chr17: 72,838,168–72,856,007), *GRIN2D* (chr19: 48,898,132–48,948,187), *GRIN3A* (chr9: 104,331,635–104,500,862). *GRIN3B* (chr19: 1,000,437–1,009,723) was sequenced only in patients. The detailed descriptions of the samples screened for each *NMDAR* gene are mentioned in Table 1. Primers were designed using the ExonPrimer program (Helmholtz Center Munich, Munich, Germany) from the UCSC genome browser (Supplementary Table 3). PCR products were sequenced at the McGill University and Genome Quebec Innovation Centre in Montreal, QC, Canada (<http://gqinnovationcenter.com>) on a 3730XL DNA Analyzer System (Applied Biosystems, Life Technologies Corporation, Carlsbad, CA, USA). In each case, variations were confirmed by re-amplifying the fragment and resequencing of the samples from the proband and available parents, using reverse and forward primers. PolyPhred (version 6.11), PolySCAN (v.3.0) and Mutation Surveyor (version 3.10; Soft Genetics, State College, PA, USA) were used for mutation detection analysis. The PolyPhen (<http://genetics.bwh.harvard.edu/pph>), SIFT (<http://blocks.fhrc.org/sift/SIFT.html>), Panther (<http://www.pantherdb.org/tools/csnpScoreForm.jsp>) and SNAP (<http://www.rostlab.org/services/SNAP/>) programs were used to predict the overall severity of the missense mutations. In all instances, default parameters were used for each program.

Results

We found three *de novo* mutations in patients with sporadic ASD or SCZ in *GRIN2A* and *GRIN2B* (Table 2). We also identified one truncating mutations in *GRIN2C*, four truncating mutations in *GRIN3A* and recurrent truncating mutations in *GRIN3B*, both in patients and controls (Table 3). By contrast, we found no truncating mutations in patients or in controls in the *GRIN1*, *GRIN2A*, *GRIN2B* and *GRIN2D* genes. All three

Table 1 Genes and samples screened

	Samples			
	SCZ	ASD	Controls QNTS	Controls
<i>GRIN1</i>	429 subjects (292 trios)	142 trios	285 trios	Not analyzed
<i>GRIN2A</i>	429 subjects (292 trios)	142 trios	Not analyzed	283 subjects
<i>GRIN2B</i>	429 subjects (292 trios)	428 subjects (404 trios)	Not analyzed	283 subjects
<i>GRIN2C</i>	429 subjects (292 trios)	428 subjects (404 trios)	285 trios	283 subjects
<i>GRIN2D</i>	429 subjects (292 trios)	142 trios	Not analyzed	283 subjects
<i>GRIN3A</i>	429 subjects (292 trios)	142 trios	285 trios	283 subjects
<i>GRIN3B</i>	429 subjects (292 trios)	428 subjects (404 trios)	Not analyzed	Not analyzed

Abbreviations: ASD, autism spectrum disorders; SCZ, schizophrenia; QNTS, Quebec Newborn Twin Study. Number of trios indicates individuals having parental DNA from both parents available.

Table 2 Details of *de novo* mutations identified in NMDAR subunits

Gene	Variant type			Sequence change			Samples				In silico prediction for amino acid substitution			
	Protein change	Chr	Chr position ^a	Exon	cDNA change	SCZ	ASD	SIFT	Polyphen	SNAP	SIFT	Polyphen	SNAP	
GRIN2A	Missense	16	9 858 499	14	c.2902G>A	1 (292 trios)	0 (142 trios)	0,15 (tolerated)	0.069 (benign)	Neutral	0,15 (tolerated)	0.069 (benign)	Neutral 89%	
	Silent	16	9 857 732	14	c.3669C>T	1 (292 trios)	0 (142 trios)	/	/	/	/	/	/	
GRIN2B	Missense	12	13 720 084	12	c.2473T>G	0 (292 trios)	1 (404 trios)	0,18 (tolerated)	1.567 (possibly damaging)	Non neutral 70%	0,18 (tolerated)	1.567 (possibly damaging)	Non neutral 70%	

Abbreviations: ASD, autism spectrum disorders; chr, chromosome; NMDAR, N-methyl-D-aspartate receptor subunit; SCZ, schizophrenia. Number of trios indicates individuals having parental DNA from both parents available.

PolyPhen, <http://genetics.bwh.harvard.edu/pph>. SIFT, <http://blocks.fhcrc.org/sift/SIFT.html>. SNAP, <http://www.roslab.org/services/SNAP>.

^aPositions according to genome build 37.

de novo mutations and truncating mutations in *GRIN2C* and *GRIN3* were reported neither in the most recent 1000 genome release nor in hapmap.

Two *de novo* mutations were identified in the *GRIN2A* gene. The first in a patient with SCZ with no familial history of SCZ was a missense (c.2902G>A; p.A968T) located in the intracellular domain of the protein. It was not predicted as having important functional consequence based on *in silico* analysis (SIFT, PolyPhen and SNAP; Table 2); nevertheless, it modified an amino acid that is highly conserved in mammals. The second one was a silent mutation, c.3669C>T identified in another patient with sporadic SCZ, and was not predicted to be damaging by *in silico* analysis. Paternal age was 39 and maternal age was 29 for this patient. These mutations were not found in the remaining 1707 chromosomes that had been sequenced. None of these patients had a mental retardation with 11 and 12 years of education, respectively.

A missense mutation (c.2473T>G; p.L825V) was identified in the *GRIN2B* gene in a patient with ASD that had no family history of neuropsychiatric disorders or ID. The mutation was confirmed by resequencing blood DNA of this patient and his parents. Paternal age was 36 and maternal age was 32 for this patient. This mutation was absent in the remaining 2279 chromosomes and was predicted by *in silico* analysis as relatively damaging, as it occurred in the last transmembrane domain that is highly conserved among species.

We further analyzed the prevalence of truncating mutations in our subjects. We did not identify any truncating mutations in the *GRIN1*, *GRIN2A*, *GRIN2B* and *GRIN2D* genes in 854 individuals including 429 patients with SCZ, 142 with ASD and 283 healthy controls (285 controls from general population for *GRIN1*). By contrast, we identified the same truncating nonsense mutation in the *GRIN2C* gene (c.54G>A; p.W18X) in two individuals. This mutation was first identified in an ASD patient, and this variant was transmitted from his normal father, whereas there was history of mild learning disabilities in the mother. We also found this mutation in a control from the general population.

We also identified four truncating mutations in the *GRIN3A* gene. Two of them were indels creating frameshift in the open-reading frame and creating nonsense codon, one in an individual from the population control cohort (c.2349delC; p.P783PfsX23) and one in a healthy control (c.2424_2425delAA; p.Q808QfsX5). The two other truncating mutations were single-nucleotide variants. The first one (c.1522C>T; p.Q508X) was identified in a patient diagnosed with catatonic SCZ and was inherited from the mother who had clinical characteristics ranging in the SCZ spectrum. The second one (c.679G>T; p.E227X) was observed in a young control from the general population. Finally, we observed frequent (10%) truncating mutations in the *GRIN3B* gene in the same proportion as described elsewhere in the general population.⁵⁶

Overall, we identified 260 exonic variants in the seven sequenced *NMDARs* genes, including 159 variants that were not referenced in the dbSNP database (Build 132; Supplementary Tables 1 and 2). These variants were all confirmed manually by genotyping the proband and in the respective proband's parents, when available, to determine their inheritance. To avoid rare variants that could be linked to

Table 3 Details of rare truncating mutations in NMDAR subunits

Gene	Sequence change						Samples		
	Variant type	Protein change	Chr	Chr Position ^a	Exon	cDNA change	ASD frequency	SCZ frequency	Controls frequency
<i>GRIN2C</i>	Nonsense	p.W18X	17	72 851 178	1	c.54G > A	1/428	0/429	1/568
<i>GRIN3A</i>	Nonsense	p.E227X	9	104 499 583	1	c.679G > T	0/142	0/429	1/568
	Nonsense	p.Q508X	9	104 433 172	3	c.1522C > T	0/142	1/429	0/568
	Indel	p.P783PfsX23	9	104 432 345	3	c.2349delC	0/142	0/429	1/568
	Indel	p.Q808QfsX6	9	104 390 612_104 390 613	4	c.2424_2425delAA	0/142	0/429	1/568

Abbreviations: ASD, autism spectrum disorders; chr, chromosome; NMDAR, N-methyl-D-aspartate receptor subunit; SCZ, schizophrenia.

^aPositions according to genome build 37.

ethnic origin, we removed variants coming from patients with non-Caucasian ethnicity. When doing so, we did not observe either accumulation of rare variants in patients compared with controls or accumulation of more damaging variants in patients, based on *in silico* analysis.

Discussion

Here, we provide the first systematic study of sequence alterations in all *NMDARs* genes in ASD and SCZ, and report several potentially damaging and/or *de novo* mutations. Indeed, we identified potentially damaging *de novo* mutation in *GRIN2A* and *GRIN2B* associated with sporadic cases of SCZ and ASD, respectively, and truncating mutations in *GRIN2C*, *GRIN3A* and *GRIN3B*, whereas neither truncating nor *de novo* mutations were found in *GRIN1* and *GRIN2D*.

Deep sequencing technologies have brought new insight in the genetics of neurodevelopmental disorders including SCZ. Recent reports of the literature have shown that there should be, in average, less than two *de novo* mutations in the coding sequence per individual and less than one *de novo* that would have an effect on amino acid sequence per individual.^{44,57,58}

We here report on three *de novo* mutations, two in *GRIN2A* in two patients suffering from SCZ, and one in *GRIN2B* in a patient with autism. The mutation in *GRIN2B* is one of the mutations quoted in a previous paper reporting an overall rate of *de novo* mutation in autism or SCZ.⁴⁴ In this former paper that considers the overall rate of *de novo* mutation, based on partial results, the mutation was unfortunately misclassified as being in a patient with SCZ, whereas it is definitely in a patient with ASD. These mutations were not inherited from unaffected parents, and the patients with *de novo* mutation in *GRIN2A* or *GRIN2B* did not have a predisposing or causative mutation in 600 other synaptic genes that have been sequenced, including SCZ-risk genes, such as *DISC1* (data not shown). Hence, it could be hypothesized that these mutations could be associated with the pathogenesis of ASD or SCZ. This is further supported by the observation that the missense mutation in *GRIN2B* is predicted by *in silico* analysis to be relatively damaging and it occurred in the last transmembrane domain that is highly conserved among species. In addition, one of the *de novo* mutations identified in the *GRIN2A* gene, is located in a domain that is highly conserved in mammals and vertebrates, and is known to be the target of post-translational modifications, especially phosphorylation. In particular, it has been recently proposed that abnormalities

in the neuregulin 1-ErbB4 pathway, which has been implicated in SCZ and known to contribute to NMDAR hypofunction, acts through NMDARs phosphorylation by the Src tyrosine kinase.^{18,19}

Several studies have already screened for mutations in some *NMDARs* genes in neurodevelopmental diseases. Williams *et al.*⁵⁹ screened for mutations in the *GRIN1* and *GRIN2* genes in patients with SCZ, but did not identify any variants that were clearly relevant to the phenotype, with only a single missense variant identified among 368 individuals (including 184 controls). Endeley *et al.*²⁵ screened for mutations and cytogenetic abnormalities in the *GRIN2A* and *GRIN2B* genes in patients with ID and found rare damaging *de novo* mutations and truncating mutations in these genes, indicating that these mutations can be pathogenic. Recently Hamdan *et al.*²⁶ screened *GRIN1* in patients with non-syndromic ID and identified a *de novo* missense and one amino acid in-frame insertion that altered the receptor's activity, the control population in this latter paper being the same as the one used in the present report. Lastly, an exome sequencing study published this year, identified several potentially causative *de novo* mutations on patients with ASD, one of them being in *GRIN2B*, within a splice site.²⁹ As we do not provide direct evidence of a functional effect of these mutations, we cannot definitely conclude on the causal effect of *de novo* mutations identified in *GRIN2A* and *GRIN2B* genes. It should however be emphasized that there were no truncating mutations in at least 854 individuals (including 283 healthy volunteers) in *GRIN1*, *GRIN2A*, *GRIN2B* and *GRIN2D*, supporting the hypothesis of a high selection pressure on these genes.

In addition to *de novo* mutations, we identified truncating mutations in the *GRIN2C*, *GRIN3A* and *GRIN3B* genes in patients, population controls and/or healthy volunteers. The rate of mutations in the *GRIN3B* gene was in the same proportion as described in the general population, with around 10% of the individuals homozygous for truncating mutation.⁵⁶ As this relatively high rate of mutation and as *GRIN3B* is mainly expressed in motor neurons, and thus a role in mental disorders as SCZ and ASD is unlikely, the gene was not further explored in controls.^{56,60}

The presence of truncating mutations in controls suggests that loss of function of these subunits leads to little or no effect. For the *GRIN2C* subunit, we identified the same truncating mutation in a patient with autism and in a control. Although the global functioning and learning abilities of this

control appear to be in the normal range, his relatively young age at the time of his evaluation precludes from reaching firm conclusions on the effect of this truncating mutation on his future development.

On the other hand, the loss of function in one allele from the *GRIN3* family genes do not appear to have dramatic consequences as truncating mutations were identified in healthy controls, suggesting some functional compensations. NMDAR containing GRIN2 subunits show large differences in their pharmacological and electrophysiological properties, including glutamate affinity, modulation by glycine, sensitivity to Mg^{2+} and channel kinetics. As they also show a distinct spatial and chronological pattern of expression, it has been suggested that these receptors may have unique roles in brain development.¹ Unlike most NMDAR that contain GRIN2 subunits, which typically require glycine and glutamate for activation, NMDAR that contain both GRIN1 and GRIN3 subunits are activated by glycine alone. When co-expressed with GRIN1 and GRIN2 subunits, GRIN3 subunits can act as negative modulators, reducing single-channel conductance and Ca^{2+} permeability.⁶¹ Thus, it is not surprising that rare damaging mutations could have differential effects in the GRIN2 and GRIN3 family of NMDARs.

We identified a large number of rare variants that are not in the dbSNP databases. Based on *in silico* analysis, these variants do not appear to be pathogenic on their own. Further, most of them are inherited from unaffected parents and some of them could be linked to the ethnic origin, as they were more frequent in the Tunisian samples. When excluding the mutations found in subjects that were not of European Caucasian origin, we did not observe any accumulation of rare or more damaging variants in patients compared with controls. Nevertheless, we cannot exclude complex gene interaction that could occur with variants in other genes.

In conclusion, our data support the hypothesis that rare *de novo* mutations in *GRIN2A* or *GRIN2B* could account for some cases of sporadic SCZ or autism. They also further support the implication of NMDARs in psychiatric disorders with neurodevelopmental origin. Nevertheless, a distinct pattern is observed between the different NMDARs. Truncating mutations in *GRIN2C* and *GRIN3* were found in patients and controls, whereas no truncating mutations were found in *GRIN1*, *GRIN2A*, *GRIN2B* and *GRIN2D*. This could indicate that *GRIN1*, *GRIN2A*, *GRIN2B* and *GRIN2D* families are crucial in normal brain development and function and/or that functional compensation could occur to counteract the loss of one allele in *GRIN2C* and *GRIN3* family genes.

Conflict of interest

The authors declare no conflict of interest.

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