

1 **Differential Functional Consequences of *GRIN2A* Mutations Associated 2 with Schizophrenia and Neurodevelopmental Disorders**

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14

15 **Abstract**

16 Human genetic studies have revealed rare missense and protein-truncating variants in
17 *GRIN2A*, encoding for the GluN2A subunit of the NMDA receptors, that confer significant
18 risk for schizophrenia (SCZ). Mutations in *GRIN2A* are also associated with epilepsy and
19 developmental delay/intellectual disability (DD/ID). However, it remains enigmatic how
20 alterations to the same protein can result in diverse clinical phenotypes. Here, we
21 performed functional characterization of human GluN1/GluN2A heteromeric NMDA
22 receptors that contain SCZ-linked GluN2A variants, and compared them to NMDA
23 receptors with GluN2A variants associated with epilepsy or DD/ID. Our findings
24 demonstrate that SCZ-associated *GRIN2A* variants were predominantly loss-of-function
25 (LoF), whereas epilepsy and DD/ID-associated variants resulted in both gain- and loss-
26 of-function phenotypes. We additionally show that M653I and S809R, LoF *GRIN2A*
27 variants associated with DD/ID, exert a dominant-negative effect when co-expressed with
28 a wild-type GluN2A, whereas E58Ter and Y698C, SCZ-linked LoF variants, and A727T,
29 an epilepsy-linked LoF variant, do not. These data offer a potential mechanism by which
30 SCZ/epilepsy and DD/ID-linked variants can cause different effects on receptor function
31 and therefore result in divergent pathological outcomes.

1 Introduction

2 NMDA (N-methyl-d-aspartate) receptors (NMDAR) play integral roles in synaptic
3 transmission, development and plasticity^{1,2}. Human genetic association studies, as well
4 as postmortem transcriptomics and proteomics studies have implicated synaptic
5 dysfunction in schizophrenia (SCZ) risk³. Additionally, several lines of evidence support
6 NMDAR/glutamate hypofunction as a mechanism underlying SCZ pathophysiology: (i)
7 NMDAR antagonists like phencyclidine and ketamine induce SCZ-like symptoms in
8 humans⁴; (ii) autoimmune-NMDAR encephalitis can present with SCZ-like symptoms⁵;
9 (iii) mouse models of NMDAR hypofunction display some phenotypic and biological
10 similarities to SCZ⁶; (iv) extensive human genetics data implicate glutamate receptor
11 signaling alterations in SCZ⁷⁻¹².

12 The recent Schizophrenia Exome Sequencing Meta-Analysis (SCHEMA) has
13 identified *GRIN2A*, which encodes the GluN2A subunit of the NMDAR, as one of ten
14 genes with exome-wide significance for SCZ risk¹³. *GRIN2A* has additionally been
15 implicated as a SCZ risk gene by genome-wide association studies^{14,15}. *GRIN2A* is highly
16 intolerant to mutations leading to a loss-of-function (LoF) phenotype in humans, implying
17 that *GRIN2A* insufficiency is highly detrimental to evolutionary fitness¹⁶. The association
18 of *GRIN2A* with SCZ is largely driven by protein truncating variants (PTVs)¹³, which are
19 predicted to be loss-of-function (LoF)¹⁷. A recent multi-omic study of *Grin2a* heterozygous
20 and homozygous null mutant mice has shown that loss of a single copy of *Grin2a* leads
21 to brain-wide transcriptomic changes and phenotypic features reminiscent of human SCZ,
22 including heightened resting gamma oscillation power in electroencephalogram
23 recordings, reduced brain activity in the prefrontal cortex, and a hyperdopaminergic state
24 in the striatum^{18,19}.

25 In addition to SCZ, *GRIN2A* is associated with epilepsy, intellectual disability (ID)
26 and developmental delay (DD)²⁰. Association with these neurodevelopmental disorders
27 appears to be mediated predominantly through missense mutations that are often
28 localized in the transmembrane and linker domains of GluN2A²⁰⁻²². Functional analyses
29 of disease-associated variants of *GRIN2A* linked to conditions such as epilepsy or DD/ID
30 have demonstrated both gain- and loss-of-function consequences²²⁻²⁶.

1 Here we present an *in vitro* functional analysis of three groups of *GRIN2A*
2 mutations: i) 11 SCZ-linked mutations found by the SCHEMA study¹³; ii) seven mutations
3 found in the control human subjects of the SCHEMA study¹³ and one mutation from the
4 Genome Aggregation Database (gnomAD)²⁷ (referred collectively hereafter as control
5 mutations); iii) four mutations associated with severe DD/ID and epilepsy^{20,22,28}. Our data
6 demonstrate that all tested early PTVs and a subset of missense variants associated with
7 SCZ display a LoF phenotype, defined as either reduced current density or an increase
8 in the glutamate EC₅₀ or both, while DD/ID-associated missense variants display either a
9 total loss of response to glutamate or a gain-of-function (GoF) effect compared to wild-
10 type receptor function. Additionally, we provide evidence that DD/ID-associated LoF
11 variants can exert a dominant-negative effect when co-expressed with wild-type *GRIN2A*,
12 whereas SCZ-associated LoF variants do not, providing a potential pathomechanistic
13 model that might provide insight into predicting phenotype severity of *GRIN2A* variants.

14 **Results**

15 **Mutant selection and construct expression in HEK cells**

16 NMDARs are heterotetrameric ligand-gated ion channels that consist of two GluN1
17 (encoded by *GRIN1*) and two glutamate-binding GluN2 subunits (encoded by *GRIN2A-D*). Each GluN subunit of NMDARs consists of four domains: an extracellular amino-
18 terminal domain (ATD), a clamshell-shaped ligand-binding domain (LBD) which is
19 composed of two non-adjacent segments of the polypeptide, known as S1 and S2, a
20 transmembrane domain (TMD) including three transmembrane helices (M1, M3, and M4)
21 and a membrane re-entrant loop (M2) joined by short linker regions, and an intracellular
22 carboxy-terminal domain (CTD) (Fig. 1A, Fig. S1). We selected a representative set of
23 SCZ-linked mutations found by the SCHEMA study¹³ that reside within different domains
24 of GluN2A including one PTV in the ATD (E58Ter), one PTV (Y700Ter) and four missense
25 mutations in the LBD including two with MPC (Missense badness, PolyPhen-2,
26 Constraint²⁹) pathogenicity score > 3 (hereafter referred as mis3; L794M, M788I), and
27 two with 2 < MPC < 3 (hereafter referred as mis2; Y698C, G784A), three missense
28 mutations in the linker/TMD regions (mis3: Q811P; mis2: I605M, G591R), and two
29 mutations in the CTD (mis: I1295T; Frameshift: L1377FS) (Fig. 1A, Fig. S1).

1 We also selected eight control mutations: seven of which were identified within a
2 control population of 97,322 individuals from the SCHEMA study¹³. Among the SCHEMA
3 study's control population, 50,437 individuals had no documented psychiatric diagnoses,
4 and were compiled from 11 global collections that had previously participated in common
5 variant association studies, and 46,885 samples were gathered as part of the gnomAD
6 consortium initiative with no association with psychiatric and neurological conditions.
7 These selected control mutations had no occurrence in SCZ cases and included four in
8 the LBD (mis2: K707R, Q671H, A727V, I775M), two in the M1-M2 linker region (mis2:
9 F576L, R586K), and one in the CTD (PTV: K1339Ter). We also selected one missense
10 mutation from the gnomAD database in the CTD, V967L, because it is the most prevalent
11 *GRIN2A* missense variant in the general population and has no clinical phenotype (Fig.
12 1A, Fig. S1).

13 Additionally, to elucidate the specific effects of SCZ-linked mutations, we
14 characterized one epilepsy-associated mutation A727T²⁸ in the LBD, and three missense
15 *GRIN2A* mutations associated with DD/ID in the linker/TMD regions: M653I, S809R, and
16 L812M^{20,30} the latter of which has previously been characterized²² (Fig. 1A, Fig. S1).

17 We transfected HEK cells with plasmids encoding for GFP-tagged human *GRIN1*
18 fused with a self-cleaving 2A peptide and wild-type or mutated human *GRIN2A*, allowing
19 us to express GluN1 and GluN2A proteins at a 1:1 ratio (Fig. 1B). We confirmed robust
20 expression of the constructs after 24h visually as the majority of cells were GFP-positive.
21 Biochemistry analysis by Western blot, showed that cells transfected with PTVs located
22 in the N-terminal half of the protein, E58Ter and Y700Ter, showed no detectable
23 expression of GluN2A at the size corresponding to wild-type GluN2A (Fig. 1C, Fig. S2).
24 In case of Y700Ter, a faint band near the expected size for the Y700Ter fragment (~ 80
25 kDa) was observed (Fig. S2). Both E58Ter and Y700Ter introduce premature termination
26 codons in their mRNA transcripts, leading to their degradation by nonsense-mediated
27 decay (NMD). NMD is a well-established cellular mechanism designed to degrade mRNA
28 transcripts containing premature termination codons or nonsense mutations¹⁷. However,
29 PTV transcripts are more likely to escape NMD when they are downstream of 50-55
30 nucleotides before the last exon-exon junction³¹, or reside in the last exon³². This could
31 explain why the PTVs nearer to the carboxy-terminal end of the protein (L1377Frameshift

1 and K1339Ter, both located in the last exon of *GRIN2A*) did not lead to decreased protein
2 expression in HEK cells. For the K1339Ter variant, an approximately fivefold increase in
3 expression was observed compared to the wild-type GluN2A ($p = 0.07$; Fig. 1D). The
4 increased expression could potentially be attributed to the functional disruption of a di-
5 leucine motif positioned at residues 1319-1320, situated in close proximity to the
6 K1339Ter mutation site. This motif has been established to play a regulatory role in the
7 endocytosis of NMDARs containing GluN2A³³. All *GRIN2A* missense variants tested,
8 except G784A, were expressed at comparable or higher levels than wild-type GluN2A
9 (Fig. 1C-D).

10

11 **Functional characterization of SCHEMA and non-SCHEMA mutations**

12 We next recorded whole-cell currents from HEK cells expressing wild-type or
13 mutant NMDARs using a high-throughput automated planar patch-clamp system
14 (Syncropatch 384PE). Cells were held at -60 mV during the recording and currents were
15 evoked using the liquid handler to “puff” increasing concentrations of glutamate (1, 3, 10,
16 30, and 100 μ M) in the constant presence of a saturating concentration of glycine (30 μ M)
17 (Fig. 2A). Of the 11 selected SCZ-linked variants, both PTVs located in the N-terminal
18 half of the protein (E58Ter, Y700Ter) as well as one mis3 (Q811P) and one mis2 (Y698C)
19 variants displayed electrophysiological characteristics that were significantly different
20 from the wild-type control (Fig. 2B-C). E58Ter and Y700Ter showed no response to any
21 concentration of glutamate (Fig. 2B), consistent with the lack of protein expression of
22 these mutants (Fig. 1C-D). The missense variant Q811P, predicted to have a high degree
23 of deleteriousness (mis3), showed a fivefold increase in glutamate EC₅₀ compared to the
24 wild-type control (Q811P: $19.91 \pm 2.11 \mu$ M; WT: $3.92 \pm 0.257 \mu$ M, $p < 0.0001$) (Fig. 2C),
25 but no change in maximal response at a saturating concentration (100 μ M) of glutamate.
26 The mis2 variant Y698C showed a fivefold reduction in maximal response to the
27 saturating concentration of glutamate compared to wild type (Y698C: $-10.40 \pm 1.47 \text{ pA/pF}$;
28 WT: $-46.88 \pm 8.67 \text{ pA/pF}$, $p < 0.0001$) (Fig. 2B) as well as a twofold increase in glutamate
29 EC₅₀ (Y698C: $6.55 \pm 0.731 \mu$ M; WT: $2.79 \pm 0.255 \mu$ M, $p < 0.05$) (Fig. 2C). The other tested
30 SCZ-linked mutations did not significantly differ in any of the measured characteristics
31 compared to the wild-type GluN2A/GluN1 NMDARs (Fig. 2D-E, Fig. S3; Table 1).

1 Together, our data demonstrate that SCZ-linked missense *GRIN2A* variants either had
2 no measurable effect on NMDAR function in our electrophysiological assay or resulted in
3 reduced glutamate potency or efficacy. Notably, the Q811P and Y698C variants
4 demonstrated a LoF phenotype, similar in direction but less severe than the SCZ-linked
5 PTVs. Consistent with our results, prior characterization of another SCZ-linked missense
6 variant identified by the SCHEMA study, A716T, revealed a LoF phenotype of increased
7 glutamate EC₅₀²⁵.

8 None of the control variants tested (seven from the SCHEMA study and one from
9 gnomAD) showed a significantly different EC₅₀ or maximal response to glutamate (Fig.
10 2D-E, Fig. S4; Table 1).

11 All four epilepsy and DD/ID-linked mutations showed significantly different
12 electrophysiological phenotypes from wild type (Fig. 2B-E, Fig. S5). A727T, associated
13 with epilepsy²⁸, showed a significant reduction in maximal response to glutamate (A727T:
14 -6.595 ± 3.13 pA/pF; WT: -29.01 ± 4.54 pA/pF, p < 0.005) (Fig. 2B, D), consistent with
15 previous characterizations²⁵. L812M displayed a fivefold reduction in glutamate EC₅₀
16 (L812M: 1.90 ± 0.266 μM; WT: 9.66 ± 1.53 μM, p < 0.0001) (Fig. 2C, E) and no change
17 in maximal current density (Fig. 2D), consistent with previous studies showing that this
18 variant enhances glutamate potency²². By contrast, both M653I and S809R did not
19 respond to any concentration of glutamate, despite protein expression levels similar to
20 wild type (Fig 1D), indicating a LoF phenotype (Fig. 2B, Fig. S5).

21 In summary, out of the 11 SCZ-linked *GRIN2A* variants characterized by this study,
22 two tested PTVs located in the N-terminal half of the protein led to complete loss of current
23 in response to glutamate and two out of the eight missense variants showed a partial LoF
24 phenotype. Of the eight characterized control variants, seven missense variants and one
25 PTV, none demonstrated any changes as compared to the wild-type receptor. Notably,
26 all tested epilepsy and DD/ID-associated missense variants were significantly different
27 from wild type and displayed either a severe LoF or a GoF phenotype (Fig. 2 D-E; Table
28 1).

29

30 **Dominant-negative phenotype as a differentiator of severe DD/ID and epilepsy/SCZ**

1 Our data demonstrate that *GRIN2A* missense variants associated with either
2 epilepsy, DD/ID or SCZ can result in LoF of the NMDAR. However, the clinical features
3 of these disorders, all of which stem from heterozygous mutation of *GRIN2A* in humans,
4 are distinct, raising the question whether there are functional differences in the *in vivo*
5 consequences of these variants. In order to better reflect the human disease state, we
6 co-expressed the SCZ, epilepsy, and DD/ID-linked variants, that were found to result in
7 LoF, with the wild-type variant and investigated their effects on NMDAR function. Having
8 previously demonstrated that the DD/ID-linked missense variant M653I produces a
9 comparable amount of protein as the wild-type receptor (Fig. 1C-D), we first tested
10 whether the null electrophysiological phenotype caused by this variant in heterologous
11 cells was a result of dysfunctional trafficking of the receptor to the cell surface. To this
12 end, we conducted surface biotinylation of HEK cells expressing either the M653I variant
13 or wild-type GluN2A. We validated the assay by also probing for insulin receptor B,
14 endogenously expressed on the surface of HEK cells, and beta-actin, endogenously
15 expressed primarily in the cytoplasm. This assay demonstrated that M653I-containing
16 NMDARs are present at the cell surface at a comparable level to the wild-type receptor
17 (Fig. 3A, Fig. S6).

18 We then tested whether either SCZ-linked or epilepsy and DD/ID-linked variants
19 would exert a dominant-negative effect on NMDAR function when co-expressed with an
20 equal amount of wild-type *GRIN2A* (as well as the obligate *GRIN1* subunit). For this
21 experiment, we included the DD/ID-linked variants, M653I and S809R, the epilepsy-linked
22 variant, A727T, as well as Y698C, a SCZ-linked missense variant, all with a LoF
23 phenotype (Fig. 2B-E), and E58Ter, an early termination mutation that produces no
24 protein (Fig. 1C) and therefore should not have a dominant-negative phenotype. No
25 difference in glutamate EC₅₀ or maximal current density was observed between the 1:1
26 WT:E58Ter co-expressing cells and the wild-type NMDAR (Fig. 3B-C). Interestingly, the
27 1:1 WT:Y698C and 1:1 WT:A727T co-expressing cells demonstrated no change in
28 glutamate EC₅₀ and their maximal responses did not exhibit a statistically significant
29 difference when compared to cells co-expressing 1:1 WT:E58Ter (Fig. 3B-C). Cells co-
30 expressing 1:1 WT:M653I and 1:1 WT:S809R *GRIN2A*, however, showed a decrease in
31 glutamate EC₅₀ and a significant reduction of ~70% and ~45% in maximal current

1 compared with cells co-expressing 1:1 WT:E58Ter, respectively (WT:M653I vs
2 WT:E58Ter: $p < 0.0001$; WT:S809R vs WT:E58Ter: $p = 0.0002$; Fig. 3B-C),
3 demonstrating that the DD/ID-linked variants, M653I and S809R, exert a dominant-
4 negative effect on NMDAR function, whereas the epilepsy-linked variant, A727T, and the
5 SCZ-linked variants, Y698C and E58Ter, do not.

6 These data uncover a potential mechanism by which similar functional
7 consequences (LoF) of GluN2A alteration can lead to different effects on receptor
8 function, and therefore divergent pathological outcomes.

9

1 Discussion

2 A growing number of disease-associated variants have been identified in *GRIN2A*,
3 highlighting the need for mechanistic data to establish connections between genetic
4 variants and the resultant pathological phenotypes. Here, we present electrophysiological
5 data for a representative set of *GRIN2A* variants associated with SCZ, epilepsy or DD/ID,
6 and propose a pathomechanistic model that could potentially aid in predicting phenotype
7 severity of *GRIN2A* variants.

8 In agreement with previous findings in *GRIN2A*, *GRIN1* and *GRIN2B*^{20,25,34,35}, our
9 data suggest that the location of mutations in the structure of GluN2A can inform
10 predictions of their functional consequences and pathogenicity. We noted that all disease-
11 associated *GRIN2A* missense variants that led to an electrophysiological phenotype
12 significantly different from wild type were located either within the LBD (Y698C and
13 A727T) or linker/TMD regions (Q811P, L812M, S809R and M653I) of GluN2A while none
14 of the mutations putatively localized in the CTD (I1295T, L1377FS, V967L, and
15 K1339Ter) demonstrated electrophysiological changes. Additionally, we show that
16 among disease-associated *GRIN2A* variants, the DD/ID mutations localized within the
17 linker/TMD region of GluN2A resulted in either a severe LoF (no current) or GoF
18 (enhanced glutamate potency) consistent with prior characterizations of mutations from
19 this regions^{20,22–25,28,36}, while SCZ/epilepsy-linked missense mutations in the LBD
20 displayed either no effect or partial LoF (reduced current or glutamate potency). These
21 data align with previous studies showing that missense variants in the linker/TMD regions
22 are associated with more severe DD/ID, while missense variants in the LBD generally
23 result in LoF and are associated with less severe abnormalities with mild to no ID^{20,25}.

24 It is noteworthy, however, that heterologous overexpression of *GRIN2A* variants in
25 HEK cells may not reveal deficits in trafficking and other post-translational mechanisms
26 resulting from these mutations. For instance, reduced glutamate potency has been shown
27 to often be associated with decreased surface expression of NMDARs in neurons²⁵. Thus,
28 it is possible that disease-associated missense mutations that lead to reduced glutamate
29 potency in our study (such as Q811P and Y698C) lead to a more severe LoF phenotype
30 when expressed in neurons due to cumulative effects on glutamate potency and receptor
31 trafficking. Additionally, it has been shown that rare variants at the interfaces between

1 subunits of NMDARs might not necessarily affect the receptor function but lead to
2 impaired receptor trafficking²⁵ resulting in a LoF phenotype. An example of such a variant
3 in our study is the SCZ-linked L794M which forms a non-bonded contact with GluN1 upon
4 NMDAR assembly. Although we did not observe any effect of L794M on NMDAR function,
5 it is plausible that its expression in neurons results in LoF. Finally, the heterologous
6 overexpression system might not reveal the functional consequences of mutations in the
7 CTD which has an indispensable role in NMDAR-mediated intracellular signaling and
8 synaptic physiology. Thus, further studies of the tested *GRIN2A* variants expressed in
9 neurons are necessary to elucidate the impact of these mutations in a physiologically
10 relevant context.

11 Our data also offers a potential mechanistic explanation for how missense *GRIN2A*
12 variants can result in the same phenotype in the homozygous state but lead to distinct
13 neurological manifestations in a heterozygous state. We show that the DD/ID-linked
14 M653I and S809R, epilepsy-linked A727T and SCZ-linked Y698C *GRIN2A* variants all
15 result in LoF, but, when co-expressed with wild-type *GRIN2A*, only M653I and S809R
16 exert a dominant negative effect. Looking into the position of these mutations in the 3D
17 structure of the GluN1/GluN2A NMDAR, we noted that the M653I and S809R mutations,
18 but not Y698C or A727T, are located at the interface between GluN2A and GluN1, and
19 introduce intramolecular interactions with GluN1, which are not present in the wild-type
20 NMDAR. Thus, it is possible that the new interactions introduced by M653I and S809R
21 mutations are rigidifying the assembly and thereby causing a negative effect on the entire
22 assembly. This is consistent with previous findings showing that mutations in the protein-
23 protein interface can cause a dominant negative effect by poisoning the assembly of the
24 protein complex^{37,38}. These data suggest that the consequences of *GRIN2A*
25 haploinsufficiency as a result of DD/ID-associated missense variants might be more
26 severe as compared to SCZ/epilepsy-associated missense variants and PTVs, despite
27 the two groups having similar effects on NMDAR function in a homozygous state. These
28 data highlight the importance of analyses of *GRIN2A* variants in both homozygous and
29 heterozygous conditions to effectively differentiate between various potential
30 mechanisms. Together, our findings offer a better understanding of the relationship
31 between genetic variant, NMDAR dysfunction, and disease phenotype, and could

- 1 potentially contribute to development of pharmacologic strategies to correct NMDAR
- 2 function.

1 Methods

2 DNA Constructs

3 The wild-type construct was synthesized by Genscript Biotech by adding eGPF
4 (accession JN204884.1), GRIN1 (accession NM_000832.7), and GRIN2A (accession
5 NM_000833.5) to the pcDNA3.1+P2a backbone. Mutant constructs were generated by
6 Genscript Biotech using site-directed mutagenesis. All constructs were transformed into
7 NEB Turbo Competent E. coli (High Efficiency) (New England Biolabs C2984H) and
8 purified using the NucleoBond Xtra Maxi EF kit (Machery Nagel 740424).

9

10 Cell Culture and Electroporation

11 HEK 293-T cells (Sigma-Aldrich 12022001) were cultured at 37 °C, 5% CO₂ in DMEM,
12 high glucose, GlutaMAX Supplement, pyruvate (Thermo 10569044) supplemented with
13 10% fetal bovine serum and 5% pen-strep (DMEM + FBS/PS). Cells were electroporated
14 using the MaxCyte STX electroporator. Briefly, 50-75% confluent cells were washed with
15 PBS, lifted, pelleted, and resuspended at a final density of 100 million cells/mL in
16 electroporation buffer (MaxCyte proprietary buffer). 100 µL of this mixture was loaded in
17 the electroporation cassette (OC-100), and electroporated using the preset “HEK 293”
18 protocol with 25 µg of the appropriate DNA construct concentrated to ≥ 5 µg/µL. For co-
19 expression in the dominant-negative experiments, cells were electroporated with a
20 mixture of 12.5 µg of both constructs. Post-transfection, 10 mg of DNase I (Stemcell
21 Technologies 07900) was added to the cells, and they were allowed to recover for 30 min
22 at 37 °C, 5% CO₂, and then rescued into DMEM + FBS/PS containing 15 µM DCKA, 8
23 µM APV, and 2 µM MK801.

24

25 Whole-Cell Recording

26 Cells were recorded using the Syncropatch 384PE 24h after electroporation in a whole-
27 cell configuration. Transfected cells were washed twice with PBS and treated with
28 Accutase (Stemcell Technologies 07920) for 5 min, and then pelleted and resuspended
29 in extracellular solution (in 8 mM glucose, 4 mM KCl, 10 mM HEPES, and 145 mM NaCl
30 (pH 7.4)) at a density of 400K cells/mL. We used a cesium-containing internal recording
31 solution to minimize the contribution of endogenous cationic current during the recording

1 (in 20 mM EGTA, 10 mM CsCl, 110 mM CsF, 10 mM HEPES, and 10 mM NaCl, pH 7.2).
2 The patch process consisted of a series of different negative pressures and negative
3 voltages to foster the giga-seal to the glass (REF PMID: 29736723). To enhance the seal
4 formation the cells were transiently exposed to a high Ca^{2+} recording solution “seal
5 enhancer solution” (in 80 mM NaCl, 8 mM glucose, 60 mM NMDG, 4 mM KCl, 10 mM
6 HEPES, and 10 mM CaCl_2 , pH 7.4). The cells were washed four times (replacing half of
7 the volume of the well) with standard recording solution (in mM 80 NaCl, 8 glucose, 60
8 NMDG, 4 KCl, 10 HEPES, and 6 CaCl_2 , and 30 μM glycine, pH 7.4) before starting the
9 recording. After gaining electrical access (whole-cell configuration), cells were held at a
10 holding potential of -60 mV, and NMDAR currents were evoked by a “puffing addition
11 protocol” (PMID: 36340694) with increasing concentrations of glutamate (0, 1, 3, 10, 30,
12 100 μM) in recording solution. Glutamate puffs exposed the cells to the glutamate-
13 containing solution for ~250 ms before being washed out with the recording solution in
14 the stack. All the glutamate was removed completely after each puff. To remove any
15 traces of glutamate, after each puff half of the volume of the well was replaced with fresh
16 recording solution in between each addition of ligand.

17

18 **Electrophysiological analyses**

19 Groups of 5 variants at a time were transfected and recorded along with a wild-type
20 construct as a reference, and the results for each of the variants were normalized to this
21 wild-type control. Different batches of cells electroporated with wild-type control showed
22 a variability in the maximal response to glutamate, but not in the glutamate EC_{50} , likely
23 related to the electroporation efficiency for each particular batch of cells. Cells were
24 selected to be analyzed based on having a seal resistance of ≥ 50 M Ohm for all ligand
25 application steps and a maximal current ≥ 50 pA. Traces were analyzed between 800 ms
26 and 1400 ms after initiation of stacked ligand addition. All current values were normalized
27 to the capacitance of the recorded cell to calculate current density. Glutamate EC_{50} values
28 were calculated for each cell using a 3-parameter agonist-response model with a Hill
29 slope of 1.0, Response = Bottom + Concentration*(Top-Bottom)/(EC50 + Concentration).

30

31 **Immunoblot analysis**

1 Cells were electroporated as described. After 24h, cells were washed 2x with PBS and
2 lysed in 1% SDS containing protease inhibitors (Sigma 4693159001) and nuclease
3 (Sigma E1014). The protein concentrations of cell lysate were determined using
4 Bicinchoninic acid assay (BCA; Pierce 23227). To equalize protein concentrations,
5 samples were diluted with 4X SDS-Sample buffer (Boston Bioproducts BP-111R; to a
6 final concentration of 1X) and water. The diluted samples were then left at room
7 temperature for 20 min. 25 µg of protein in an equal volume were loaded for each sample
8 on 3-8% Tris-acetate polyacrylamide gels; the gels were run using Tris-acetate SDS
9 running buffer at constant voltage. Proteins were transferred to 0.2 µm Nitrocellulose
10 membranes using semi-dry transfer (BioRad Transblot Turbo; 25V 30 min). Membranes
11 were blocked using 5% milk in Tris-buffered saline supplemented with 0.1% Tween-20
12 (TBST) for 1.5h at room-temperature (RT). Membranes were then probed overnight at 4
13 °C with primary antibody (Novus rabbit anti-NMDAR2A NB300-105, 1:2000 or CST rabbit
14 anti- Insulin Receptor β 4B8, 1:1000) in 1% milk TBST with gentle rotation. After three 10-
15 min washes in TBST, membranes were incubated with 1:5,000 HRP-conjugated (Jackson
16 Immunoresearch) anti-IgG antibody in 1% milk TBST for 60 min at RT. All membranes
17 were then washed three times in TBST and imaged on the ChemiDoc MP (BioRad)
18 platforms. Membranes were then re-probed with an HRP-conjugated mouse anti-beta
19 actin antibody (Sigma A3854) for 1h at RT in 5% milk TBST and imaged again as
20 described.

21

22 **Surface Biotinylation**

23 Cells were electroporated as described before. After 24h, cells were washed two times
24 with PBS supplemented with 1 mM CaCl₂ and 1 mM MgCl₂. Cells were then incubated in
25 1 mg/mL sulfo-NHS-LC biotin in PBS + Mg/Ca (Thermo 21335) on a flat surface for 30
26 min-2h at 4 °C. Cells were then quenched in 20 mM Tris in PBS + Mg/Ca for 15 min at
27 4°C. Cells were then lysed in 20 mM Tris.Cl (pH 7.5), 150 mM NaCl, 0.2% Triton-X 100,
28 1% SDS in water supplemented with protease inhibitors and nuclease. After quantification
29 of lysate with BCA, 100 µg of lysate was removed and diluted to 0.2% SDS by water, and
30 rotated at 4 °C for 30 min. 80 µL of neutravidin-agarose beads (Thermo 29204) was added
31 to this lysate and the solution was rotated overnight at 4 °C. Afterwards, the solution was

1 spun down, and a sample of the flowthrough was taken for analysis, and the remaining
2 beads were washed in 200 μ L lysis buffer three times. Beads were then eluted with 4X
3 SDS-Sample buffer.

4

5 **Statistics**

6 All quantitative data is given as mean \pm SEM, unless stated differently. All
7 statistical comparisons were done using one-way Brown-Forsythe and Welch ANOVAs
8 with Dunnett's T3 multiple comparison correction.

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7

8 **Author contributions**

9 Z.F. and M.S. designed the study with assistance from D.B.N. and J.Q.P.. N.S., N.B. and E.K.
10 performed all the experiments with assistance from D.B.N.. N.S., D.B.N., and Z.F. performed data
11 analysis. S.I. and A.J.C. carried out the protein structure analysis. N.S. and Z.F. wrote the
12 manuscript with inputs from all co-authors.

13 **Data availability statement**

14 All data generated or analyzed during this study are included in this published article (and its
15 Supplementary Information files), and are available from the corresponding author on reasonable
16 request.

17 **Additional Information**

18 M.S. is cofounder and SAB member of Neumora Therapeutics, and serves on the SAB of Biogen,
19 ArcLight, Vanqua Bio, and Proximity Therapeutics.

20

1 **Figure Legends**

2 **Figure 1. GRIN2A mutation selection, construct design, and validation of expression in** 3 **HEK cells.**

4 (A) All pathogenic and non-pathogenic variants selected for characterization in this study, mapped
5 on the domain structure of GluN2A. Missense variants from SCZ cases were colored according
6 to the predicted impact (MPC score) in the function. PTV, protein-truncating variant; DD/ID,
7 developmental delay/intellectual disability; FS, frameshift.

8 (B) Diagram of the construct transfected into HEK 293T cells for the functional characterization
9 of *GRIN2A* variants. The plasmid was designed with a P2A sequence between the two genes to
10 control the expression of both transcripts with one high efficiency promoter (CMV), and to assure
11 equimolar protein production of GFP-tagged *GRIN1* and wild-type or mutated *GRIN2A* for
12 electrophysiological characterization.

13 (C) Western blots probing for GluN2A and β-ACTIN in lysates of HEK cells transiently transfected
14 with *GRIN1*-*GRIN2A* constructs to express wild-type or mutant NMDARs. The blots presented
15 here are cropped, and the original blots are presented in Supplementary Figure 2.

16 (D) Quantification of GluN2A expression by Western blot. All values are normalized to wild-type
17 GluN2A expression. Data are shown as mean + SD; n = 3 for each
18 GluN2A variant. Statistical significance was assessed using Brown-Forsythe and Welch ANOVA
19 with Dunnett's T3 multiple comparisons test, no conditions were found to be significantly different
20 from wild type.

21 22 **Figure 2. Schizophrenia and DD/ID-associated GRIN2A mutations demonstrate both gain-** 23 **and loss-of-function effects.**

24 (A) Whole-cell recording protocol using Syncropatch to record NMDAR currents from HEK cells
25 transiently transfected with the construct shown in Fig.1B. Cells were puffed with different
26 concentrations of glutamate stacked in between different volumes of reference solution (light gray
27 triangles, inset on the right panel). Each puff results in a ~250 ms transient of glutamate exposure
28 to the receptors. After each glutamate application half of the volume of the well was replaced with
29 fresh reference solution ("Reference exchange", gray bars) to minimize desensitization due to
30 remnant glutamate in the well. Glu, glutamate; ref, reference solution.

31 (B) Averaged current traces of wild-type and selected mutant NMDARs evoked by ~250 ms
32 transients of glutamate exposure of increasing concentration, in the presence of 30 μM glycine.
33 The traces were colored according to the different concentrations of glutamate.

1 (C) Averaged current density in response to increasing concentrations of glutamate in the
2 constant presence of 30 μ M glycine, normalized to maximal response, for wild-type and selected
3 mutant NMDARs. The lines indicate a nonlinear regression three-parameter fit to each dataset.

4 (D) Peak current density in response to 100 μ M glutamate, normalized to wild type's response,
5 for each mutant NMDAR.

6 (E) Glutamate EC₅₀ normalized to wild type EC₅₀, for each mutant NMDAR.

7 In **C-E** data are displayed as mean \pm SEM, n = 10-77; see Table 1 for number of cells recorded
8 per variant; statistical significance was assessed using Brown-Forsythe and Welch ANOVA with
9 Dunnett's T3 multiple comparisons test. *: p < 0.05, **: p < 0.001, ***: p < 0.0001.

10

11 **Figure 3. GRIN2A mutations associated with DD/ID, but not epilepsy or SCZ, demonstrate**
12 **a dominant-negative effect.**

13 (A) Western blot probing for GluN2A, insulin receptor beta, and β -ACTIN in the input, flowthrough,
14 and elution samples of surface biotinylation experiment done on HEK cells transiently transfected
15 with *GRIN1-GRIN2A* constructs to express wild-type or M653I mutant NMDARs. Input,
16 flowthrough, and elution represent total, internal, and surface expression respectively. IR-B:
17 insulin receptor beta. The blots presented here are cropped, and the original blots are presented
18 in Supplementary Figure 6.

19 (B) Glutamate EC₅₀ normalized to wild type EC₅₀, for each mutant NMDAR.

20 (C) Peak current density in response to 100 μ M glutamate, normalized to wild type's response, is
21 plotted for each mutant NMDAR.

22 In **B** and **C** data are displayed as mean \pm SEM; n = 80 (WT); 113 (1:1 WT:E58Ter); 27 (Y698C);
23 45 (1:1 Y698C:WT); 8 (A727T); 44 (1:1 A727T:WT); 74 (M653I); 39 (1:1 WT:M653I); 35 (S809R);
24 46 (1:1 WT:S809R) cells; statistical significance was assessed using Brown-Forsythe and Welch
25 ANOVA with Dunnett's T3 multiple comparisons test. *: p < 0.05, **: p < 0.001, ***: p < 0.0001.

26

27 **Table 1. Electrophysiological and expression data for all characterized GRIN2A mutations.**

28 Data are given as mean \pm SEM except for protein expression values which are given as mean
29 \pm SD. Statistical significance was assessed using Brown-Forsythe and Welch ANOVA with
30 Dunnett's T3 multiple comparisons test. Bolded text indicates significantly different values from
31 wild type. *: p < 0.05, **: p < 0.01, **: p < 0.001, ***: p < 0.0001. The cell colors in the table
32 indicate the pathological status of the variants. The lightest gray shade represents the SCZ-linked
33 mutation, the second light gray corresponds to control mutations, and the darkest gray shade is
34 indicative of mutations linked to epilepsy and developmental delay/intellectual disability (DD/ID).

1 **Supplemental Figure Legends**

2 **Figure S1. GRIN2A mutations mapped on human NMDAR structure.** Protein structure of
3 human NMDAR (PDB ID 6IRH) with two GluN1 (light yellow) and two GluN2A (light green)
4 subunits. GRIN2A mutations associated with SCZ (red) or epilepsy/DD/ID (blue) as well as control
5 mutations (dark gray) are mapped on the two GluN2A subunits of human NMDAR. Two SCZ-
6 associated missense mutations (Q811P, G591R), one control mutation (R586K), one DD/ID-
7 linked mutation (L812M) as well as mutations within the CTD were not mapped due to no structure
8 coverage.

9

10 **Figure S2. Uncropped western blot images of GluN2A variants.** Western blots probing for
11 GluN2A and β -ACTIN in lysates of HEK cells transiently transfected with GRIN1-GRIN2A
12 constructs to express wild-type or mutant NMDARs. The arrow in Y700Ter lane indicates a faint
13 band near the expected size for the Y700Ter fragment (~ 80 kDa).

14

15 **Figure S3. Effect of SCZ-associated GRIN2A variants on NMDAR function.** Averaged current
16 density in response to increasing concentrations of glutamate in the constant presence of 30 μ M
17 glycine, normalized to maximal response, for NMDARs containing the wild-type or SCZ-
18 associated GluN2A variants. The lines indicate a nonlinear regression three-parameter fit to each
19 dataset. Data are displayed as mean \pm SEM, n = 31-81; see Table 1 for number of cells recorded
20 per variant.

21

22 **Figure S4. Effect of non-pathogenic GRIN2A variants on NMDAR function.** Averaged current
23 density in response to increasing concentrations of glutamate in the constant presence of 30 μ M
24 glycine, normalized to maximal response, for NMDARs containing the wild-type or non-pathogenic
25 GluN2A variants. The lines indicate a nonlinear regression three-parameter fit to each dataset.
26 Data are displayed as mean \pm SEM, n = 17-70; see Table 1 for number of cells recorded per
27 variant.

28

29 **Figure S5. Effect of Epilepsy and DD/ID-associated GRIN2A variants on NMDAR function.**
30 Averaged current density in response to increasing concentrations of glutamate in the constant
31 presence of 30 μ M glycine, normalized to maximal response, for NMDARs containing the wild-
32 type or Epilepsy and DD/ID-linked GluN2A variants. The lines indicate a nonlinear regression

1 three-parameter fit to each dataset. Data are displayed as mean \pm SEM, n = 10-77; see Table 1
2 for number of cells recorded per variant.

3

4 **Figure S6. Original western blot images of the surface biotinylation experiment.** Western
5 blot probing for GluN2A, insulin receptor beta, and β -ACTIN in the input, flowthrough, and elution
6 samples of surface biotinylation experiment done on HEK cells transiently transfected with
7 *GRIN1-GRIN2A* constructs to express wild-type or M653I mutant NMDARs. Input, flowthrough,
8 and elution represent total, internal, and surface expression respectively. IR-B: insulin receptor
9 beta.

10

Figure 1

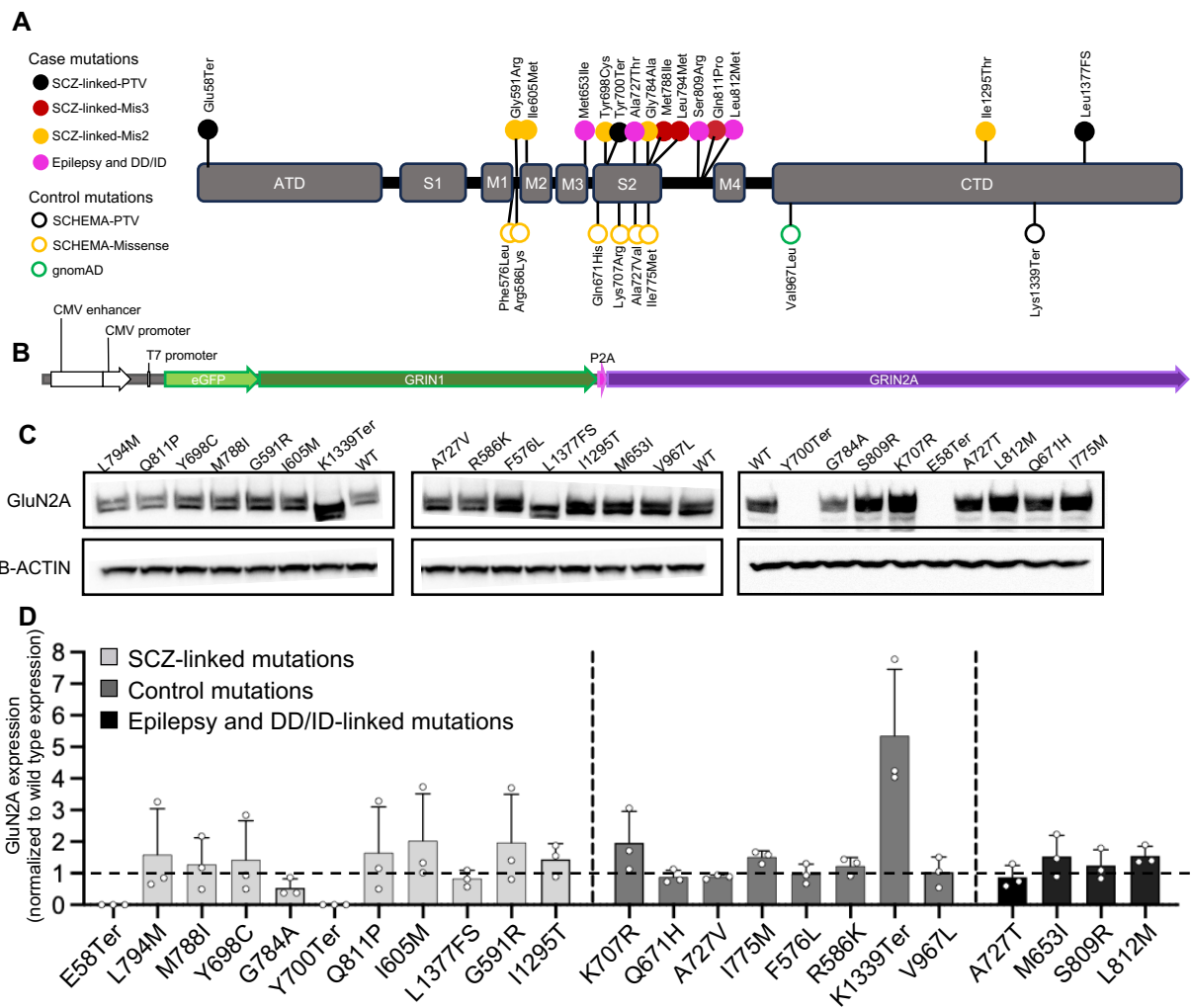


Figure 2

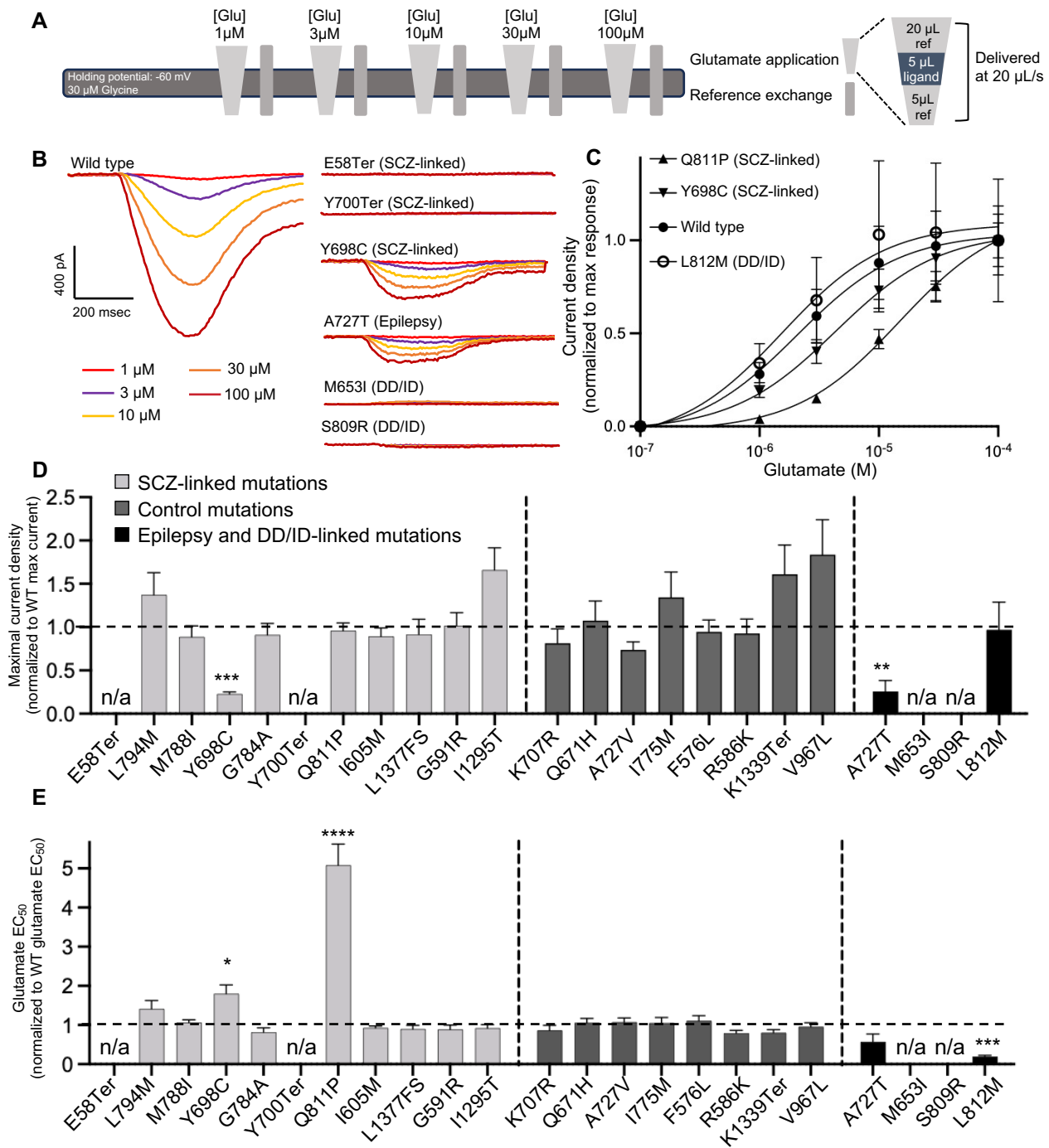


Figure 3

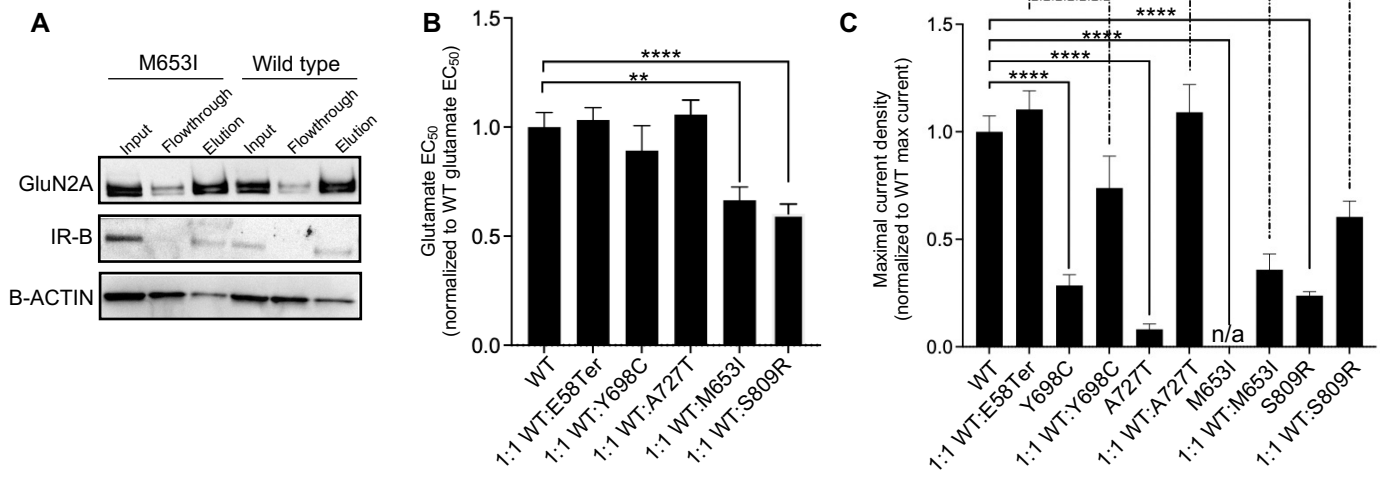


Table 1

| Variant | Location on GluN2A | Mutation Class | Glutamate EC ₅₀ (normalized to WT) | Max Amplitude (normalized to WT) | Protein Expression (normalized to WT) | Number of Cells |
|----------|--------------------|----------------|---|----------------------------------|---------------------------------------|-----------------|
| E58Ter | ATD | PTV | N/A | N/A | N/A | 41 |
| L794M | LBD (S2) | mis3 | 1.41 ± 0.21 | 1.37 ± 0.26 | 1.59 ± 0.84 | 37 |
| M788I | LBD (S2) | mis3 | 1.06 ± 0.08 | 0.89 ± 0.13 | 1.28 ± 0.84 | 71 |
| G784A | LBD (S2) | mis2 | 0.82 ± 0.11 | 0.91 ± 0.13 | 0.54 ± 0.16 | 33 |
| Y698C | LBD (S2) | mis2 | 1.80 ± 0.23 (*) | 0.23 ± 0.02 (****) | 1.42 ± 0.72 | 47 |
| Y700Ter | LBD (S2) | PTV | N/A | N/A | N/A | 36 |
| Q811P | Linker (S2-M4) | mis3 | 5.08 ± 0.54 (****) | 0.96 ± 0.09 | 1.65 ± 0.84 | 50 |
| I605M | TMD (M2) | mis2 | 0.93 ± 0.05 | 0.89 ± 0.10 | 2.03 ± 0.86 | 81 |
| L1377FS | CTD | PTV | 0.90 ± 0.09 | 0.92 ± 0.18 | 0.83 ± 0.15 | 31 |
| G591R | Linker (M1-M2) | mis2 | 0.89 ± 0.11 | 1.01 ± 0.15 | 1.97 ± 0.88 | 58 |
| I1295T | CTD | mis2 | 0.92 ± 0.09 | 1.66 ± 0.25 | 1.44 ± 0.29 | 37 |
| K707R | LBD (S2) | mis2 | 0.87 ± 0.13 | 0.81 ± 0.17 | 1.97 ± 0.57 | 18 |
| Q671H | LBD (S2) | mis2 | 1.06 ± 0.11 | 1.07 ± 0.23 | 0.88 ± 0.12 | 17 |
| A727V | LBD (S2) | mis2 | 1.07 ± 0.11 | 0.74 ± 0.09 | 0.88 ± 0.04 | 70 |
| I775M | LBD (S2) | mis2 | 1.05 ± 0.14 | 1.35 ± 0.29 | 1.51 ± 0.11 | 36 |
| F576L | Linker (M1-M2) | mis2 | 1.11 ± 0.13 | 0.95 ± 0.14 | 0.97 ± 0.18 | 35 |
| R586K | Linker (M1-M2) | mis2 | 0.79 ± 0.07 | 0.93 ± 0.17 | 1.23 ± 0.16 | 38 |
| V967L | CTD | mis2 | 0.81 ± 0.07 | 1.84 ± 0.40 | 1.03 ± 0.28 | 38 |
| K1339Ter | CTD | PTV | 0.96 ± 0.10 | 1.61 ± 0.34 | 5.35 ± 1.21 | 33 |
| A727T | LBD (S2) | mis2 | 0.57 ± 0.20 | 0.26 ± 0.13 (**) | 0.89 ± 0.21 | 10 |
| L812M | Linker (S2-M4) | mis3 | 0.20 ± 0.03 (**) | 0.97 ± 0.32 | 1.55 ± 0.17 | 13 |
| S809R | Linker (S2-M4) | mis3 | N/A | N/A | 1.24 ± 0.29 | 30 |
| M653I | TMD (M3) | mis3 | N/A | N/A | 1.53 ± 0.38 | 77 |

| |
|-------------------------------------|
| SCZ-linked mutations |
| Control mutations |
| Epilepsy and DD/ID-linked mutations |

Figure S1

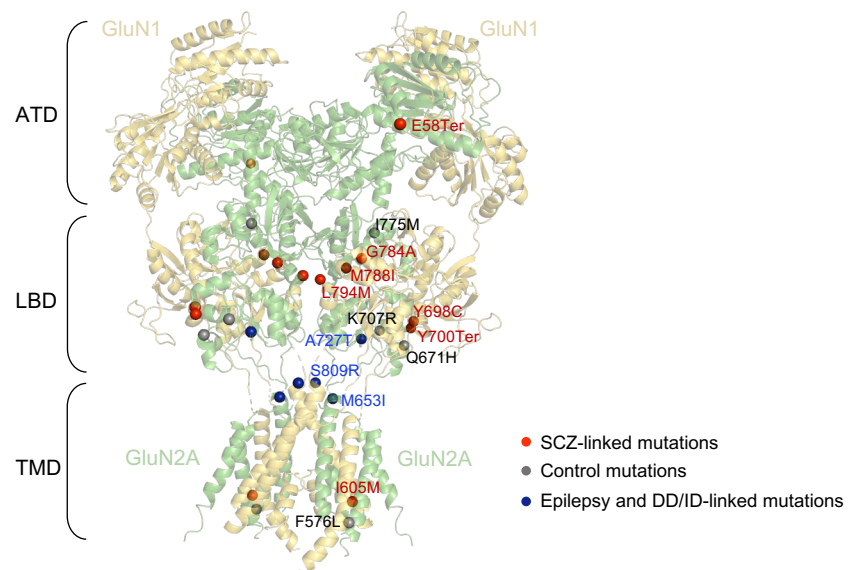


Figure S2

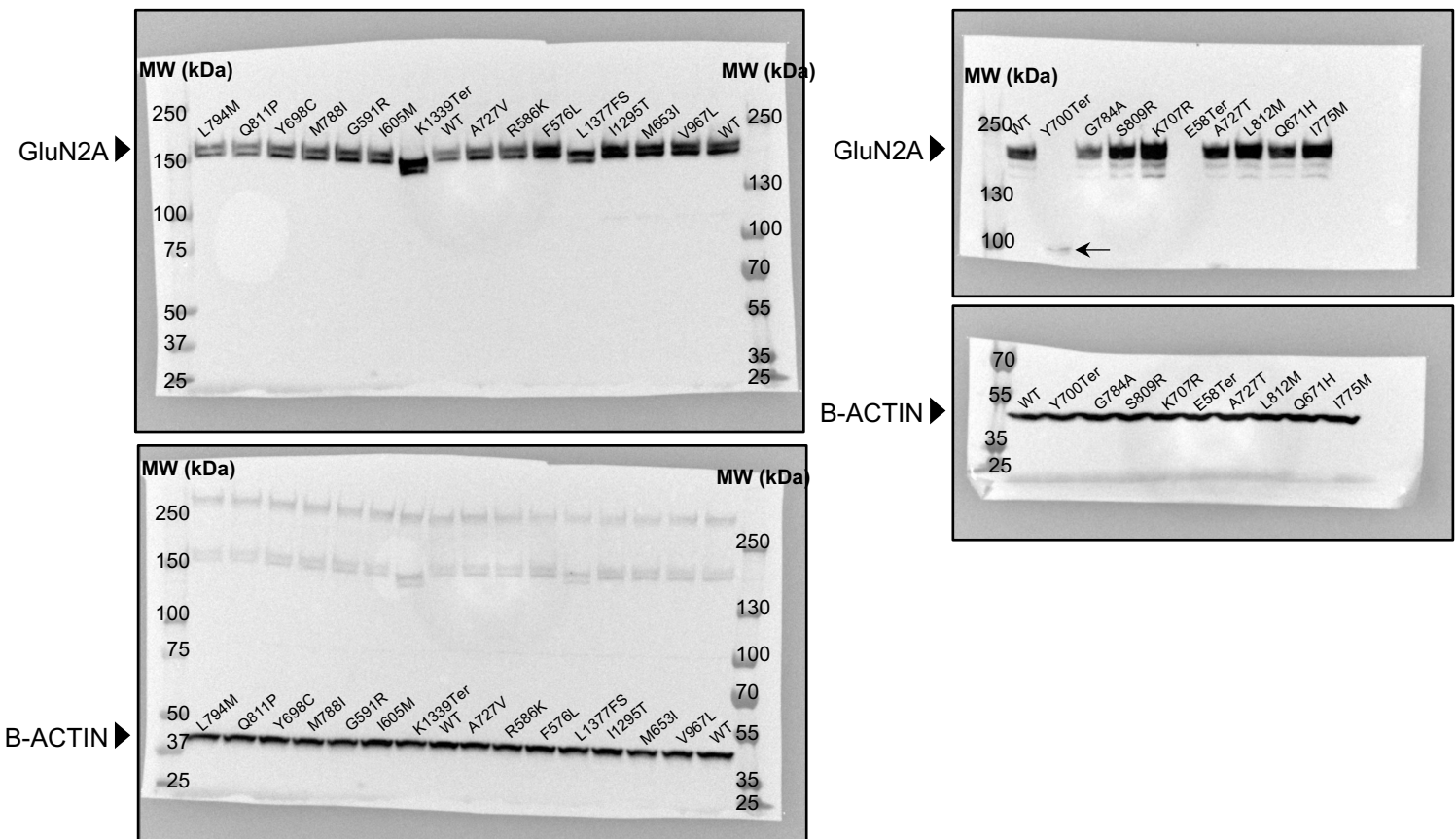


Figure S3

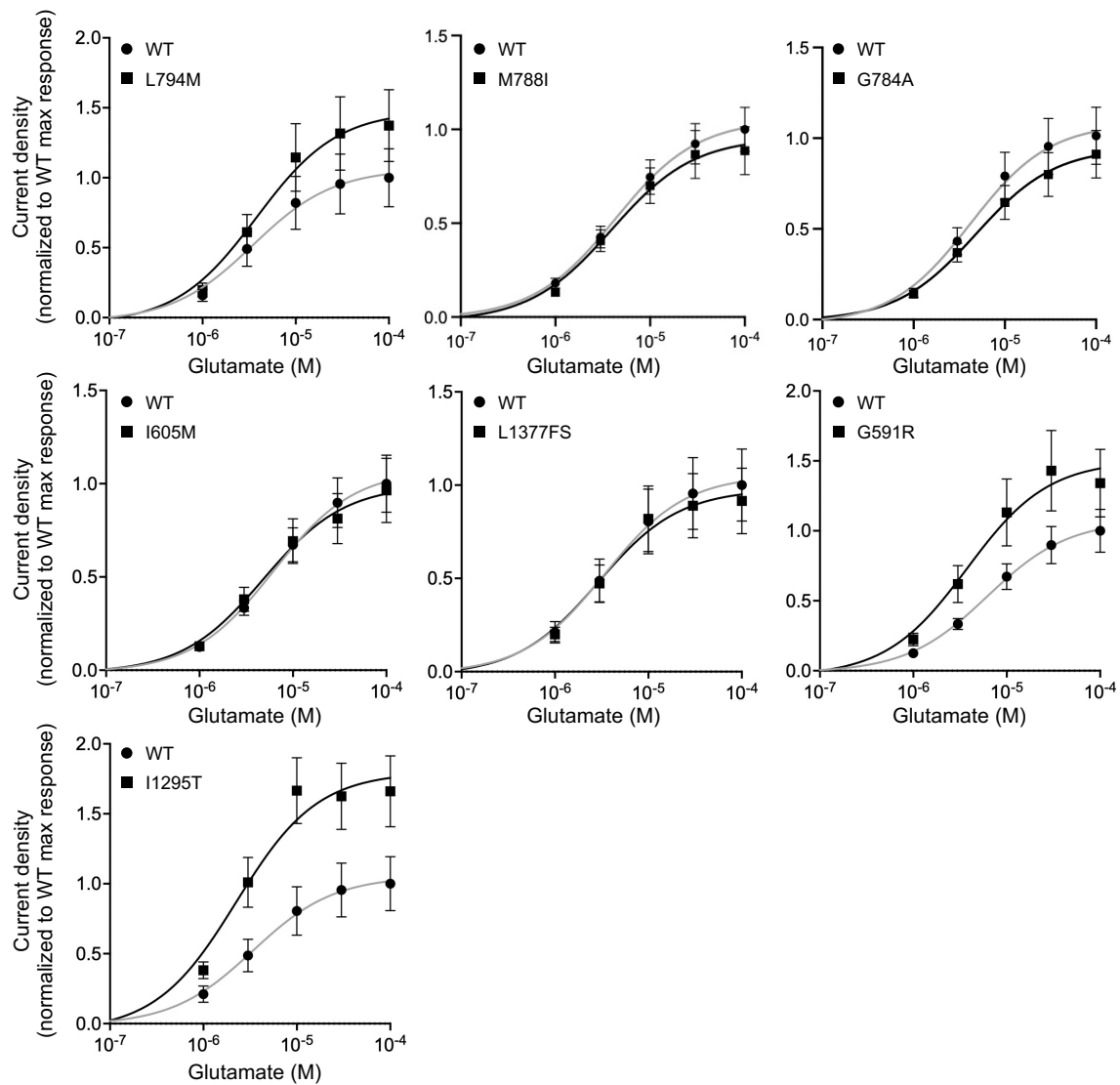


Figure S4

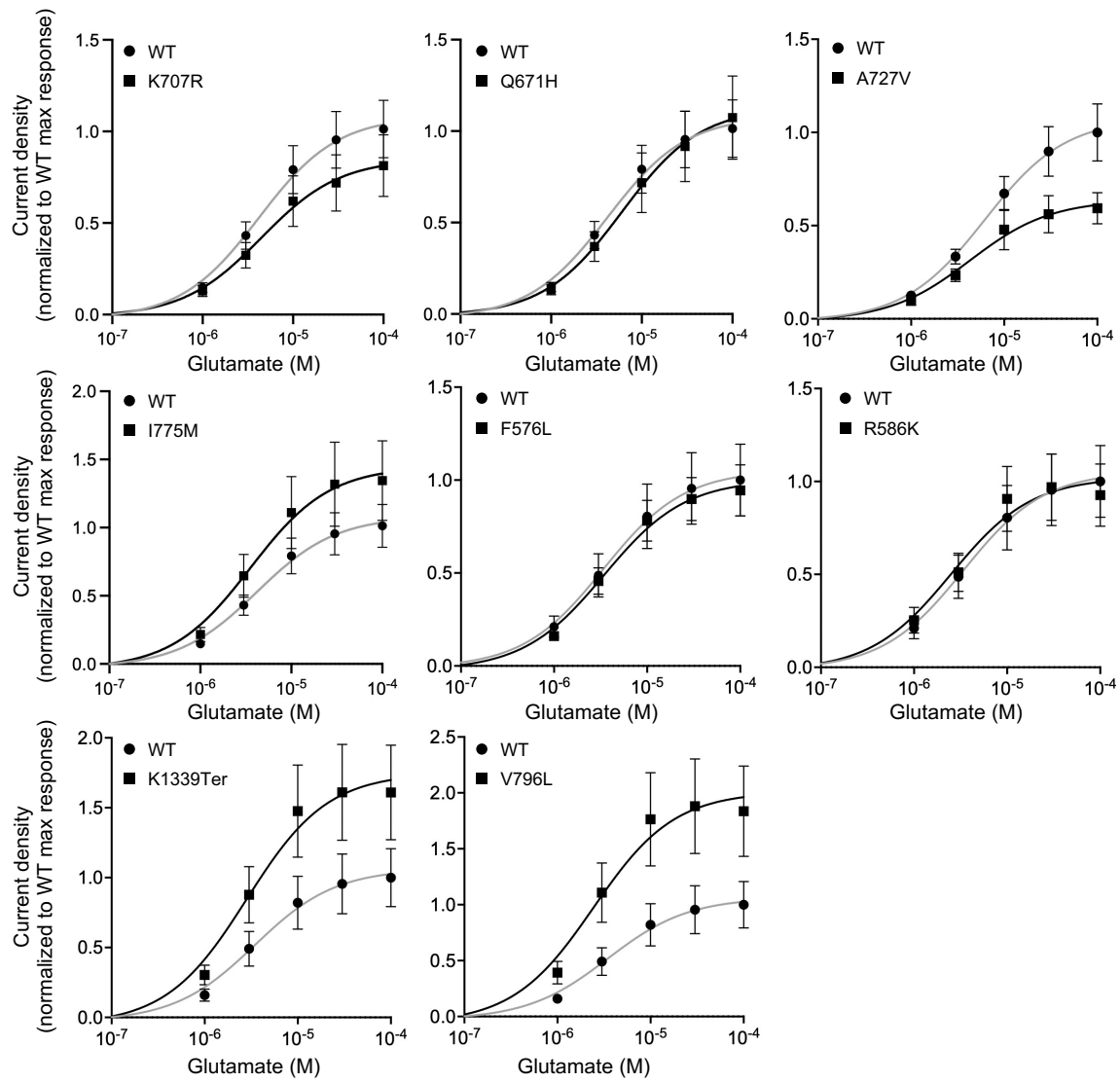


Figure S5

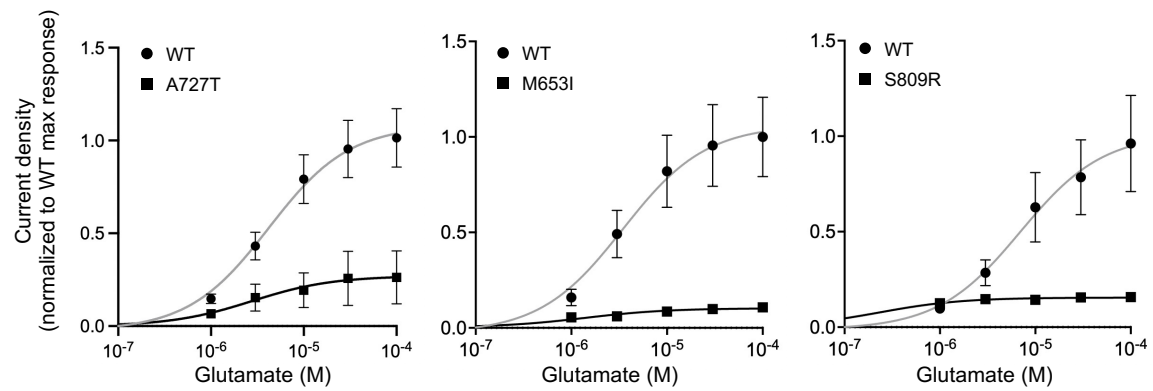


Figure S6

