NMDAR-mediated transcriptional control of gene expression in the specification of interneuron subtype identity

Vivek Mahadevan^a, Apratim Mitra^b, Yajun Zhang^c, Areg Peltekian^a, Ramesh Chittajallu^a,
 Caraoline Esnault^{b,1}, Dragan Maric^d, Christopher Rhodes^c, Kenneth A. Pelkey^a, Ryan
 Dale^b, Timothy J. Petros^c, Chris J. McBain^{a,*}

⁷ ^aSection on Cellular and Synaptic Physiology, Eunice Kennedy Shriver National Institute of Child Health
 ⁸ and Human Development (NICHD), Bethesda, 20892, MD, USA

^bBioinformatics and Scientific Programming Core, NICHD, Bethesda, 20892, MD, USA

^cUnit on Cellular and Molecular Neurodevelopment, NICHD, Bethesda, 20892, MD, USA

^dFlow and Imaging Cytometry Core Facility, National Institute of Neurological Disorders and Stroke (NINDS), Bethesda, 20852, MD, USA

13 Abstract

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Medial ganglionic eminence (MGE)-derived parvalbumin (PV)+, somatostatin (SST)+ and Neurogliaform (NGFC)-type cortical and hippocampal interneurons, have distinct molecular, anatomical and physiological properties. However, the molecular mechanisms regulating their diversity remain poorly understood. Here, via single-cell transcriptomics, we show that the obligate NMDA-type glutamate receptor (NMDAR) subunit gene *Grin1* mediates subtype-specific transcriptional regulation of gene expression in MGE-derived interneurons, leading to altered subtype identities. Notably, MGE-specific conditional *Grin1* loss results in a systemic downregulation of diverse transcriptional, synaptogenic and membrane excitability regulatory programs. These widespread gene expression abnormalities mirror aberrations that are typically associated with neurodevelopmental disorders, particularly schizophrenia. Our study hence provides a road map for the systematic examination of NMDAR signaling in interneuron subtypes, revealing potential MGE-specific genetic targets that could instruct future therapies of psychiatric disorders.

¹⁴ Keywords: Interneurons, Medial ganglionic eminence, PV, SST, Neurogliaform, NMDA

- ¹⁵ receptor, Transcriptional regulation, Neurodevelopmental disorders, Schizophrenia,
- ¹⁶ NMDA-hypofunction, Hippocampus, Frontal Cortex, Mouse model, scRNAseq

Email addresses:

vivek.mahadevan@nih.gov (Vivek Mahadevan), ryan.dale@nih.gov (Ryan Dale), tim.petros@nih.gov (Timothy J. Petros), mcbainc@mail.nih.gov (Chris J. McBain) *Corresponding author.

17 Introduction

Medial ganglionic eminence (MGE)-derived forebrain GABAergic interneurons com-18 prise the parvalbumin-containing (PV) and somatostatin-containing (SST) subpopulations 19 throughout the entire forebrain accounting for approximately 60% of all cortical interneu-20 rons [1, 2]. In addition, approximately half of all hippocampal neurogliaform-type cells 21 (NGFCs), the so called Ivy cells, originate from the MGE [3, 4]. Interestingly, though only 22 rarely found in rodent neocortex such MGE-derived NGFCs are significantly more popu-23 lous in primate neocortex, including humans [5]. While PV neurons exert robust somatic, 24 and proximal dendritic inhibition, the SST and NGFCs mediate domain-specific dendritic 25 inhibition on their downstream pyramidal neuron targets [6]. Collectively these classes of 26 interneurons shape diverse aspects of cortical and hippocampal circuit maturation during 27 development, and critically regulate information processing in mature circuits by maintain-28 ing appropriate excitation-inhibition (E-I) balance [7]. Recent evidence indicates a critical 29 role for activity, particularly through ionotropic glutamate receptors (iGluRs), in driving 30 the morpho-physiological maturation of MGE-derived interneurons [8–12]. Unlike mature 31 interneurons where iGluRs differentially contribute towards synaptic transmission, imma-32 ture and migrating interneurons express different glutamate receptor subunits including the 33 NMDA-type iGluR (NMDAR) and AMPA/Kainate-type iGluR (AMPAR/KAR) [13–15] 34 prior to the expression of any functional synapses. This becomes particularly important 35 as the developing brain contains higher ambient glutamate levels than the adult brain [16]. 36 Collectively, higher ambient glutamate, developmental expression of iGluRs and recruitment 37 of glutamatergic signaling is considered to be trophic [8, 17, 18] and thought to engage mech-38 anisms to regulate various aspects of interneuron development including morphological and 39 electrical maturation to promote appropriate circuit integration [9, 11, 14, 16, 19–22]. 40

Interneuron-specific impairments are increasingly considered central to the etiology of 41 multiple neurodevelopmental and circuit disorders [23]. The importance of interneuron-42 expressed iGluRs is most notable in psychiatric disorders exhibiting impaired NMDAR-43 associated systems [24, 25]. In the adult brain, acute pharmacological NMDAR blockade re-44 sults in circuit disinhibition and psychotic symptoms [26], mediated in-part, by the enhanced 45 sensitivity of interneuronal NMDARs to their antagonists [27]. Indeed, direct blockade of in-46 terneuron activity also precipitates distinct behavioral deficits relevant to schizophrenia [28]. 47 In particular, ablation of the obligate NMDAR subunit gene Grin1 in interneuron-specific 48 early postnatal mouse [29], but not PV-specific [30], or glutamatergic neuron-specific Grin1 49 ablation [31], resembles global Grin1-mutants [32] in their constellation of schizophrenia-like 50 behavioral aberrations. This indicates that *Grin1* dysfunction across multiple interneuron-51 subtypes precipitates schizophrenia-like abnormalities [33]. In addition, this interneuron-52 specific NMDAR-hypofunction model is sensitive to developmental age, since adult-onset 53 Grin1 loss does not result in the same phenotypes [29]. Despite the importance of develop-54 mental NMDAR function in interneurons, and its relevance to human neurodevelopmental 55 disorders, a comprehensive interrogation of the impact of developmental NMDAR ablation 56 in interneurons, particularly across MGE-derived interneurons, is lacking. 57

It is clear that during the developmental window between embryonic day (ED) 13.5 and

postnatal day (PD) ~10 [34], a combination of innate genetic programs, external environ-59 ment, and neuronal activity shapes interneuron subtype specification leading to remark-60 able diversity [2, 21, 35, 36] The NMDAR signaling complex comprises an essential node 61 for regulating gene expression via excitation-transcription (E-T) coupling in mature cir-62 cuits [37–39]. Moreover, different NMDAR subunits are widely expressed in the developing 63 brain [40] where they provide a critical source of Ca^{2+} -entry via trophic glutamate signaling 64 prior to synaptogenesis [15, 19, 41]. However, it is not clear whether the NMDAR-mediated 65 Ca^{2+} cascades in nascent and developing MGE-derived interneurons engage transcriptional 66 programs necessary for MGE-derived interneuron diversity. To investigate this, we con-67 ditionally deleted *Grin1* in MGE progenitors that give rise to cortical and hippocampal 68 PV, SST, and NGFC subsets, using the Nkx2-1-Cre mouse line [3, 4, 42]. In this model, 69 Nkx2-1-driven Cre expression is reported in cycling/proliferating MGE cells, well before the 70 cells become postmitotic, allowing for assessment of the developmental impact of embry-71 onic loss of *Grin1* activity across all subsets of MGE-derived interneurons. Applying high-72 throughput single-cell RNA sequencing (scRNAseq), we establish that NMDAR-mediated 73 transcriptional cascades promote MGE subtype identity, by regulating the expression of di-74 verse transcriptional, synaptogenic and membrane excitability genetic programs. Notably, 75 we identify numerous disease-relevant genes that are misexpressed in MGE-derived interneu-76 rons upon *Grin1*-ablation, providing a broad road map for examination of MGE-subtype 77 78 specific regulation via NMDAR signaling.

79 **Results**

sc RNA seq recapitulates cardinal MGE subtypes and a continuum of

81 molecular profiles

To examine the molecular heterogeneity of MGE-derived GABAergic interneurons 82 by scRNAseq, we microdissected frontal cortex (CX) and hippocampus (HPC) from 83 fresh brain slices obtained from PD18-20 Nkx2.1-Cre:Ai14 mouse (Figure1A, Figure1-84 Supplement1A). Ai14-TdTomato (TdT^+) single-cell suspensions were harvested by 85 fluorescence-activated cell sorting (FACS) using stringent gating constraints including via-86 bility and doublet discrimination (Figure1-Supplement1B) as previously described [43– 87 45], and subsequently processed through the 10X Genomics Chromium controller. 9064 and 88 9964 TdT⁺ cells were recovered from cortex and hippocampus respectively across 3 biolog-89 ical replicates. To minimize the effect of excitotoxicity and stress-related transcriptional 90 noise, the tissue processing, FACS, and sample collection steps were performed in buffers 91 supplemented with Tetrodotoxin (TTX), DL -2-Amino-5-phosphonopentanoic acid (APV) 92 and Actinomycin-D (Act-D) [46]. Because we observed concordant cell clustering across the 93 replicates during preliminary analysis by Seurat v3 [47, 48] (Figure1-Supplement2A), 94 the replicates were pooled for in-depth analysis. Subsequent clustering and marker gene 95 analyses revealed that ~ 62% and 33% of the TdT⁺ MGE-sorts from cortex and hippocam-96 pus respectively, express classical GABA markers including Gad1 / Gad2, Lhx6; and the 97 MGE-subclass markers *Pvalb*, *Sst*, and *Lamp5*, marking PV and SST, NGFC subsets respec-98 tively (Figure1B, Figure1-Supplement3A). While we did not recover cells expressing 99

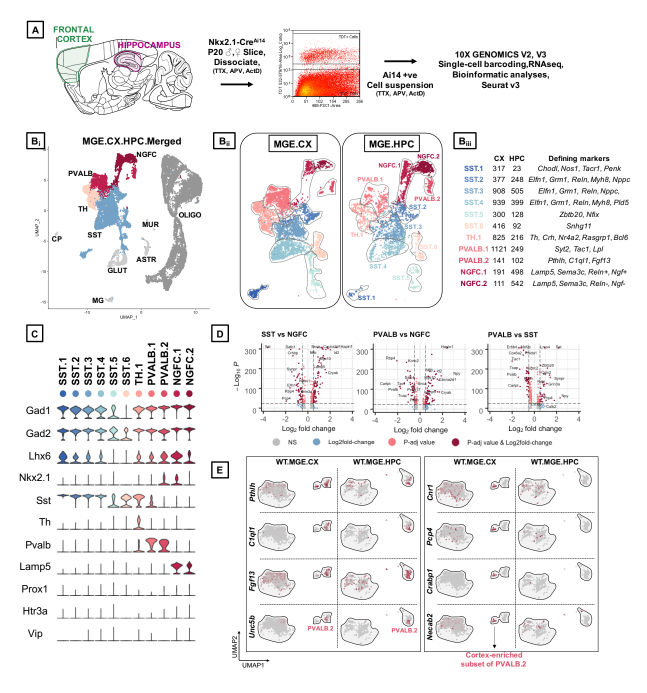


Figure 1: Identification of MGE-derived interneuron subtypes in the cortex and hippocampus. **A**, Overview of the experimental workflow. \mathbf{B}_i , Uniform Manifold Approximation and Projection (UMAP) dimensional reduction of 19,028 single-cell transcriptomes (9,064 from frontal cortex and 9,964 from hippocampus of 6 mouse brains), showing the cardinal MGE populations. Cell clusters were color coded and annotated *post hoc* based on their transcriptional profile identities (Cell type abbreviations: PVALB, Parvalbumin; NGFC, Neurogliaform; TH, Tyrosine Hydroxylase; SST, Somatostatin; GLUT, Glutamatergic; CP, Choroid Plexus; MG, Microglia; ASTR, Astrocyte; MUR, Mural; OLIGO, Oligodendrocyte). \mathbf{B}_{ii} , UMAP visualization of 11 MGE-derived interneuron subtypes from cortex (MGE.CX) and hippocampus (MGE.HPC), and the recovery of cell numbers from the subtypes. \mathbf{B}_{iii} , Table indicating the number of $Gad1/Gad2^+$ cells recovered in each MGE subtype from the cortex and hippocampus, and the defining genes enriched in each subtype. **C**, Violin plot showing the distribution of expression levels of well-known representative cell-type-enriched marker genes across the11 MGE subtypes. **D**, $-\log_{10}$ False Discovery Rate (FDR) versus log2 fold change (FC) between each of the MGE cardinal class, representing the top enriched markers at a fold change ≥ 0.5 and FDR <10e-25. **E**, UMAP representation of PVALB clusters highlighting the cortex-specific enrichment of *Pthlh*-expressing PVALB.2 subtype that is not observed in the hippocampus.

the CGE-markers *Prox1*, *Htr3a* or *Vip*, we recovered a minor fraction of cells corresponding to glutamatergic neurons, astrocytes and microglia. In addition, ~25% and 71% TdT⁺ MGEsorts from cortex and hippocampus respectively were enriched in oligodendrocytes marked by *Olig1* expression across all replicates (Figure1-Supplement 2B,2C). However, we focused our subsequent analyses on the 5656 and 3002 *Gad1* / *Gad2* positive cortical and hippocampal MGE-derived interneurons.

Unbiased cell clustering by Seurat v3 identified six subtypes of SST, two subtypes of PV, 106 two subtypes of NGFCs, and one subtype of Tyrosine hydroxylse (TH) expressing interneu-107 rons, expressing the markers Sst, Pvalb, Lamp5 and Th respectively, across the two brain 108 regions examined (Figure1C). Notably, all but two subtypes (SST.5 and SST.6) expressed 109 high levels of *Lhx6*, and 2 clusters corresponding to PV.2 and NGFC.1 expressed *Nkx2.1* 110 at this developmental time. While the PV- SST- and NGFC- clusters clearly exhibited ro-111 bust gene expression differences among each other (Figure1D), the TH cluster appeared 112 to express genes that correspond to both PV: SST clusters, including Sst and Pvalb expres-113 sions (Figure1C, Figure1-Supplement3B). Particularly, at this developmental window 114 we could not observe robustly different gene expression variances between the cortical and 115 hippocampal counterparts, barring a few marginal, but significant differences (Figure1-116 **Supplement4B**). This gave us sufficient rationale to perform subsequent analyses using 117 the MGE-derived interneurons pooled from cortex and hippocampus. 118

Among the **SST** sub clusters, SST.1-5 uniquely expresses *Chodl*, *Iqf2bp3*, *Cdh7*, 119 *Pld5* and *Nfix* respectively, while SST.6 expresses only markers that are common with 120 other SST clusters (Figure1-Supplement3B). With the exception of SST.6 the remain-121 ing SST-expressing subclusters are described in previous scRNAseq assays (Figure1-122 **Supplement5A**). For example, the *Chodl*-expressing SST.1 cluster co-express high *Nos1*, 123 Tacr1, Penk, and Npy, and it has been previously described as putative GABAergic long-124 range projections neurons [49, 50]. Clusters SST.2/3/4 express Elfn1, Reln and Grm1 125 characteristic of putative cortical martinotti and their hippocampal counterpart, oriens-126 lacunosum/moleculare (O-LM) [43, 51, 52] (Figure1-Supplement5B). Lastly, Zbtb20-127 expressing SST.5 is predicted to be septal-projecting interneurons [43]. Among the **PV sub** 128 clusters, while both PVALB.1&2 coexpresses several common markers including *Pvalb*, 129 Kcnip2, Tcap and Kcnc1 there are several notable differences between the two clusters. 130 PVALB.1 appears to contain continuous, but non-overlapping populations expressing Syt2131 representing putative fast-spiking basket cells or Rbp4/Sst containing putative bistrati-132 fied cells [1, 43, 53] (Figure1-Supplement5D) . PVALB.2 contains cells that uniquely 133 expresses Pthlh, C1ql1, Fqf13 and Unc5b representing putative axo-axonic chandelier 134 cells [43, 49, 54]. We also observed a TH cluster, which, in addition to expressing several 135 genes common to the SST: PV clusters, expresses several unique genes including Rasgrp1, 136 Bcl6, Myo1b that segregated into mutually exclusive cluster space expressing Crh or Nr4a2137 (Figure1-Supplement3B, Figure1-Supplement5B). This cluster is also described pre-138 viously as putative bistratified-like cells [43, 53]. Among the NGFC sub clusters, while 139 both NGFC.1&2 coexpress several common markers including Lamp5, Hapln1, Cacna2d1, 140 Sema3c and Id2, the NGFC.1 cluster uniquely expresses several genes like Reln, Ngf, Egfr, 141 Gabra5 that are not expressed by NGFC.2. (Figure1-Supplement3B). While the Reln+ 142

represents MGE-derived neurogliaforms, the *Reln*-population may represent putative ivy cells [43] (Figure1-Supplement5C).

While the majority of the UMAP space aligns well between the cortical and hippocam-145 pal MGE-derived interneurons, we observed some regional differences as well. (Figure1-146 **Supplement3A**_i \mathbf{A}_{iii}). (i) First, we observed an increase in the HPC-expressed NGFC.1&2 147 in comparison to their cortical counterparts, consistent with preferential localization of 148 MGE-derived NGFCs to HPC over CX in rodents [1, 3, 4]. (ii) Next, the *Pthlh*-expressing 149 PVALB.2 subcluster splits into two islands, only in the cortex and distinctly lacking from 150 the hippocampus. Only one of the PVALB.2 islands expresses C1ql1, while the other cortex-151 enriched island expresses unique markers Etv1, Cnr1, Pcp4, Crabp1, Necab2, Epha4, Crabp1 152 and Hapln1 (Figure1E, Figure1-Supplement5D). Whether this represents a novel sub-153 class of chandelier cells remains to be determined. (iii) Lasty, we also observed a distinc-154 tion in the hippocampal SST.3 corresponding to a subset of O-LM interneurons (Figure1-155 **Supplement3A**_{*ii*}). The overall MGE cell numbers indicate that the SST cells account for 156 the majority of MGE cell population recovered in the scRNAseq assay from both brain re-157 gions (Figure1-Supplement3 A_{ii} , A_{iii}). The PV and TH clusters accounted for a greater 158 share of MGE-derived interneurons in the CX than in the HPC. While it is plausible these 159 relative cell proportions may be skewed by differential survivability of these subtypes during 160 tissue dissociation, sorting and single-cell barcoding, these relative percentages were similar 161 across biological replicates. 162

¹⁶³ NMDAR signaling maintains MGE identities and subtype diversity

Because neuronal activity and glutamatergic signaling are known to regulate multiple 164 facets of interneuronal development [2, 21] [11, 36, 55], we hypothesized that the key ob-165 ligate subunit *Grin1* and the NMDAR signaling complex may play an instructive role in 166 determining MGE subtype identities. To test whether NMDAR signaling impact the devel-167 opment and function of MGE-derived interneurons, we ablated them in MGE progenitors by 168 crossing floxed-Grin1 mice with the Nkx2.1-Cre mouse line [42]. The Earliest expressions of 169 Nkx2.1 and Grin1 in the developing rodent brains occur around ~embryonic day (ED) 10.5 170 and ~ED14 respectively [56–58]. Moreover, NMDAR-mediated Ca²⁺ signaling in migrating 171 interneurons are reported by ~ED16 [15]. Because the expression and activity of Nkx2.1 pre-172 cedes Grin1 expression, we rationalized that utilizing Nkx2.1-Cre mouse will ablate Grin1 173 and NMDAR signaling in MGE progenitors from the earliest developmental point. We 174 sorted TdT⁺cells from the cortex and hippocampus of Nkx2.1-Cre: $Grin1^{fl/fl}$:Ai14 mice and 175 performed scRNAseq using the 10X platform. The scRNAseq experiments were performed 176 using juvenile mice (PD18-20) of both sexes and from the same litters as the wildtypes (WT) 177 to enable subsequent direct comparison. Similar to the WT-datasets, the MGE- $Grin1^{fl/fl}$ 178 mutants also revealed an enrichment of TdT⁺ oligodendrocytes (Figure2-Supplement3B), 179 however, we again focused our attention on the Gad1/2 positive interneurons. 180

We next performed integrated analyses of the MGE-*Grin1^{wt}* and MGE-*Grin1^{fl/fl}* cortical and hippocampal scRNAseq datasets. Applying similar unbiased clustering parameters used for the MGE-*Grin1^{wt}* analyses, we observed a total of twelve *Gad1/2* positive clusters in the integrated dataset (Figure2A, B). As a robust control, *Grin1* appeared to be absent or vastly reduced in all MGE subsets in both brain regions from MGE-*Grin1^{fl/fl}* (Figure2C),

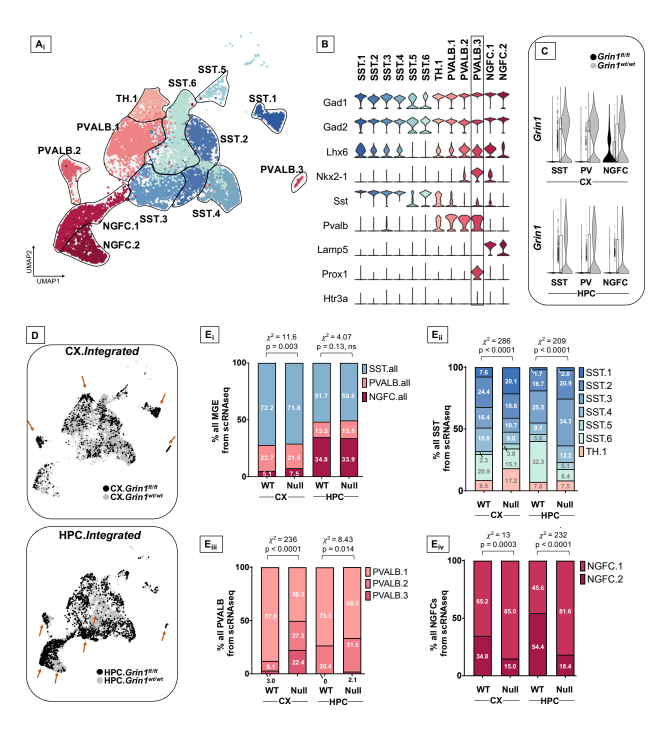


Figure 2: Altered interneuron subtype proportions upon Grin1-ablation

A, Integrated UMAP visualization of 12 subtypes of MGE-derived interneurons obtained from cortex (CX) and hippocampus (HPC) of $Grin1^{wt/wt}$ and $Grin1^{fl/fl}$ mice **B**, Violin plot showing the distribution of expression levels of well-known representative cell-type-enriched marker genes across the 12 interneuron subtypes. **C**, Violin plot from both genotypes indicating the expression of Grin1 in the cardinal of MGE-derived interneuron subtypes. **D**, UMAP representation colored by brain-region, highlighting the differential enrichments of cells (brown arrows) within interneuron subsets in Grin1-WT and Grin1-null from CX and HPC. **E**, Stacked-barplots representing the proportions of recovered cell numbers within **E**_i, pooled cardinal MGE subtypes, **E**_{ii}, SST subtypes; **E**_{iii}, PVALB subtypes and **E**_{iv}, NGFC subtypes in Grin1-WT and Grin1-WT and Grin1-WT and Grin1-WT and Grin1-WT and Grin1-NUT and Grin

but not in the *Slc17a7* expressing glutamatergic neurons (Figure2-Supplement3A). Over-186 laying the WT and NULL datasets from the brain regions revealed differential enrichments 187 among the recovered cells between the genotypes (Figure 2D). Intriguingly, Grin1-ablation 188 did not seem to alter the SST or PV recovery percentages, with the exception of a mod-180 est increase in the cortical NGFCs ($\chi^2 = 11.6$, p = 0.003), but not hippocampal NGFCs 190 $(\chi^2 = 4.07, p = 0.13)$ (Figure2E_i, Figure2-Supplement2A_i,B). However, we observed a 191 marked change in the recovery percentages of the subsets of SST, PV and NGFCs from both 192 cortex and hippocampus (Figure $2E_{ii-iv}$, Supplement $2A_{ii}$, C). Particularly, we observed 193 a robust increase in the *Chodl*-expressing cortical SST.1 population, and a decrease in hip-194 pocampal SST.6 population in MGE- $Grin1^{fl/fl}$ (CX, HPC: $\chi^2 = 286, 209$; p-value = 2.2e-16 195 for both regions). In addition, we found a reduction in the cortical PVALB.1 population, 196 and a compensatory increase in PVALB.2/3 populations in MGE- $Grin1^{fl/fl}$ (CX, HPC: χ^2 197 = 236, 8.4; p-value = 2.2e-16, 0.14). Finally, we observed an increase in the NGFC.1 along 198 with a compensatory decrease in NGFC.2 in both cortex and hippocampus (CX, HPC: χ^2 199 = 13, 232; p-value = 0.0003, 0.14).200

To independently examine whether *Grin1* ablation promotes changes in interneuron 201 abundances, we conducted immunostaining experiments to probe total TdT⁺ MGE-derived 202 interneurons and the PV / SST subtypes. First, we observed no significant change in total 203 TdT⁺ hippocampal MGE-derived interneuron density between postnatal day (PD) 30-210 204 (Figure 2-Supplement $1A_i$) similar to what was indicated in the scRNAseq cell recoveries 205 (Figure 2E_i). Next, we observed a modest decrease in cortical TdT⁺ numbers at PD30, 206 which became progressively greater by PD210 (Figure 2-Supplement $1B_i$), indicating dif-207 ferent effects of *Grin1*-ablation on total MGE cell numbers in cortex and hippocampus. 208 We observed no change in hippocampal expressed total PV/SST cell type counts at PD30 209 (Figure2-Supplement1 A_{ii}), but we noted a modest reduction in cortical PV cell type 210 counts along with an increase in cortical SST cell type counts at the same age Figure2-211 Supplement $1Bi_i$). 212

Among the differentially enriched subclusters, *Pthlh*-expressing PVALB.3 is quite no-213 table (Figure2B, Figure2-Supplement3A,B). This cortex-enriched cluster lacking in 214 the hippocampus was identified within the PVALB.2 putative-chandelier cells in the MGE-215 $Grin1^{wt/wt}$ (Figure1E, Figure1-Supplement5D), However, subsequent to integration of 216 the MGE- $Grin1^{fl/fl}$ scRNAseq dataset, it segregated as a unique cluster, far from other 217 PVALB clusters in the UMAP space. We observed *Prox1* expression in PVALB.3, which 218 is uncharacteristic of MGE-derived interneurons, additional to robust expressions of genes 219 associated with NGFCs such as *Hapln1* and *Reln* (Figure2-Supplement3A,B). More-220 over, we observed an increase in recovery of the cortical PVALB.3 cell numbers, including 221 the emergence of these cells in the hippocampus subsequent to Grin1-ablation (Figure2-222 **Supplement2A,B**). It is unclear whether the changes in marker expression reflect a true 223 change in cell identity or whether this is reflective of alterations in relative interneuron 224 subtype proportions. Nevertheless, these data demonstrate clear changes in MGE-derived 225 interneuron subtype diversity following loss early embryonic loss of *Grin1* function. 226

NMDAR signaling shapes the transcriptional landscape in MGE-derived interneurons

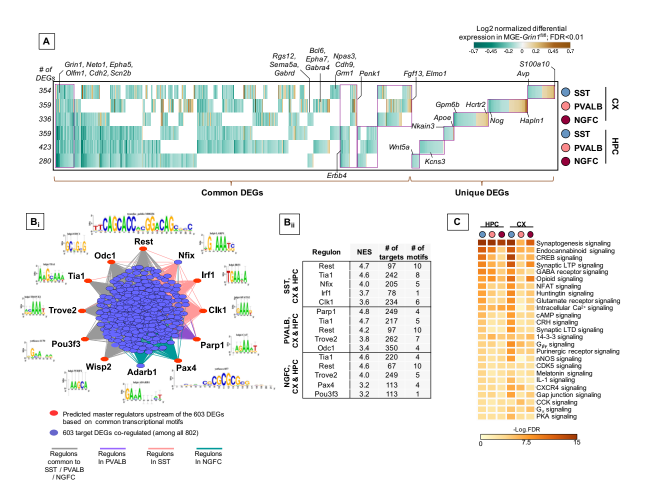


Figure 3: Cell-autonomous transcriptional changes subsequent to MGE-specific developmental *Grin1*-ablation A, Combined heatmap representing the 802 differentially expressed (DEGs) in the cortical and hippocampal MGE-derived interneurons upon *Grin1*-ablation, at a FDR<0.01 and FC>10%, as determined by MAST analysis (see details in Methods). \mathbf{B}_i , iRegulon *in silico* analysis identifying high-confidence master upstream transcriptional regulators(indicated in red) of the DEGs (indicated in lavender). Representative DNA-binding motifs are indicated next to the transcriptional regulators. \mathbf{B}_{ii} , Top five transcriptional regulators predicted by iRegulon, associated normalized enrichment score (NES) and number of predicted targets and motifs associated with each transcription factor cluster, indicated for the three interneuron subtypes (CX and HPC pooled). **C**, Ingenuity Pathway Analysis of significantly overrepresented molecular pathways in each MGE-subtype.

It is now well-established that transcriptional signatures defines the subtype identities 229 of GABAergic interneurons [59]. To examine the full range of transcriptional impairments 230 triggered by *Grin1* ablation in MGE-derived interneurons, we next performed differential 231 gene expression testing by pooling the SST / PVALB / NGFC subtypes into their cardinal 232 MGE classes to identify the genes that are differentially expressed between the genotypes. 233 For instance SST1-7 and TH.1 are pooled together as SST; PVALB1-3 are pooled together 234 as PVALB, and NGFC1-2 are pooled together as NGFC cardinal classes for this assay. At 235 a stringent false-discovery rate (FDR) <0.01, 802 genes passed the 10%-foldchange (FC) 236

threshold across the MGE subtypes from both brain regions (Figure 3A, Supplemental 237 Several interesting features were observed in the differentially expressed gene Table1). 238 (DEG) pattern upon MGE-specific Grin1-ablation. (i) Among all DEGs only $\sim 10\%$ and 1%239 are upregulated in the cortex and hippocampus respectively, while the remaining genes were 240 all downregulated (Figure 3-Supplement $1A_{ii}, B, C$). (ii) While *Grin1* ablation resulted in 241 several unique DEGs between the MGE classes, ~10 and 27% of the DEGs are common 242 within cortex and hippocampus respectively (Figure 3A, Figure 3-Supplement 2). For 243 instance, while S100a10, Hapln1, Hcrt2 are uniquely upregulated in cortical SST, PV and 244 NGFC respectively (Figure 3A), Apoe, Kcns3, Wnt5a were uniquely altered in hippocampal 245 SST, PV and NGFC respectively. In contrast, *Grin1* ablation induced common changes in 246 *Penk1* and *Erbb4* expression patterns across all MGE-derived interneuron classes in the 247 cortex and hippocampus respectively. (iii) $\sim 27-43\%$ of all DEGs were shared by MGE 248 classes across brain regions (Figure3-Supplement2A_{ii}). For example, Npas3, Cdh9, 249 Grm1 are commonly downregulated in all SST subclasses; Bcl6, Epha7, Gabra4 common 250 to PV class; and Rgs12, Gabrad, Sema5a common to NGFCs from both brain regions. (iv) 251 Lastly, 28 genes are commonly differentially expressed across both brain regions, across all 252 MGE subtypes. For example, Grin1, Neto1, Cdh2, Scn2b are commonly downregulated 253 across the board, while *Epha5*, *Olfm1* are commonly downregulated across all, but cortical 254 PV cells (Figure3A). 255

Gene expression co-regulation is intrinsic to cellular diversity [60, 61]. Since the ma-256 jority of DEGs are downregulated across the MGE subtypes, we examined whether they 257 correspond to clusters of coordinated co-regulation. We applied the iRegulon in silico frame-258 work [62], which identifies transcription factor binding motifs that are enriched in genomic 259 regions of the DEGs upon *Grin1*-ablation, and predicts the transcription factors that bind 260 to the motifs. This *in silico* analysis predicted 51 significantly enriched motifs (normalized 261 enrichment score > 3) that clustered into 10 groups by similarity, 33 of which were asso-262 ciated with transcription factors (Figure 3B, Supplemental Table 2). Put together, 10 263 transcription factors were predicted to bind with the motifs with high confidence, strongly 264 supporting targeted co-regulation of 617 among the 802 DEG genes upon Grin1-ablation. 265 Notably, the RE1-silencing transcription factor (Rest) is a master transcriptional repressor 266 that mediates the transcriptional accessibility for several synaptic genes [63], including NM-267 DAR subunits themselves [64]. It is intriguing to observe that the downregulation of the 268 DEGs upon MGE-specific Grin1-ablation are, in part, predicted to occur via Rest-mediated 269 transcriptional repression. 270

To examine the broad biological impact of the DEGs, we performed Gene Ontology 271 (GO) analyses. Broad GO analyses on all DEGs indicates that these genes serve to regu-272 late multiple molecular functions in interneurons, including regulation of GABAergic and 273 glutamatergic synapses, additional to biological pathways related to addiction and circadian 274 entrainment (Figure 3-Supplement 2B). Further classification of DEGs based on their cel-275 lular functions within the MGE subtypes revealed genes critical for regulation of membrane 276 excitability, gene expression, synaptic partnering and assembly, as well as major intracellu-277 lar Ca²⁺ signaling cascades and second messengers (Figure 3C, Figure 3-Supplement 2C, 278 Supplemental Table3). 279

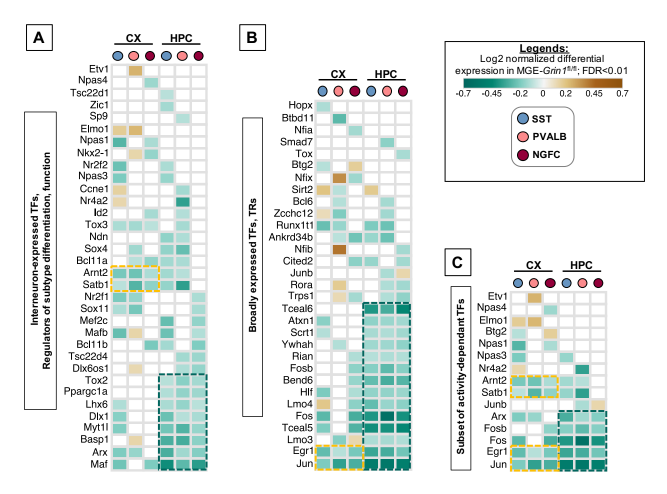


Figure 4: *Grin1*-signaling in MGE-derived interneurons are highly dedicated to the transcriptional control of interneuron identity Heatmap of log2 FC of significant DEGs in cortical and hippocampal MGE cardinal subtypes, showing a subset of \mathbf{A}_i , Transcription factors (TFs) that are previously established to regulate MGE subtype identity and function; \mathbf{B}_i , broadly expressed TFs and transcriptional regulators (TRs) that are not currently known to regulate MGE function; \mathbf{C}_i , neuronal activity-regulated TFs. Clusters of commonly differentially expressed genes in cortex and hippocampus are indicated in yellow or green boxes.

Transcription factor expression is a key component of NMDAR-mediated MGE regulation

Because transcriptional regulation underlies numerous fundamental processes including 282 the expression of other classes of genes, we next examined the DE-transcriptional regulators 283 in detail. We first examined the 67 genes that are differentially expressed upon Grin1-284 ablation and are known to mediate transcriptional regulation of gene expression. Of these, 285 35 genes are previously established to be expressed in different GABAergic interneuron 286 classes including some notable MGE-expressed transcription factors (Figure4A_i). The re-287 maining 32 are broadly expressed TFs (Figure4B_i), that include a small subset of 15 genes 288 that are regulated by neuronal activity (Figure $4C_i$). Barring a few genes, we observed 289 the majority of TFs to be down regulated in both brain regions. Intracellular Ca^{2+} sig-290

naling cascades and second messenger systems are key mediators of NMDAR signaling to 291 the nucleus for transcriptional regulation. Theoretically, an early first wave impairment of 292 Ca²⁺ signaling in *Grin1*-lacking MGE progenitors could result in transcriptional silencing 293 of the mediators of Ca²⁺ signaling cascades and second messenger systems, which would 294 sustain the transcriptional impairments. Indeed, we also observed a downregulation of var-295 ious Ca^{2+} homeostasis-regulators, kinases / phosphatases and second messengers that are 296 activated downstream of *Grin1* (Figure4-Supplement1A,B,C). Furthermore, we noted 297 that hippocampal MGE neurons had a greater proportion of DE-TFs and kinase signaling 298 cascade effectors that were downregulated across all 3 subtypes compared to their corti-299 cal counterparts. Together, this suggests that hippocampal MGE-derived interneurons may 300 be more vulnerable than cortical MGE-derived interneurons towards Grin1-mediated Ca^{2+} 301 transcriptional silencing at this age. 302

Interestingly, among the early TF cascades in the progenitors that sequentially deter-303 mine and maintain MGE fate, several members appear to be expressed at ~P20, and starkly 304 downregulated upon Grin1-ablation. For instance, Lhx6, Maf, Arx, Myt11, Dlx1 are among 305 the genes broadly downregulated across all hippocampal MGE subtypes and within spe-306 cific class(es) in their cortical parallels (**Figure4A**_{ii}). Other MGE fate-determining TFs, 307 Nkx2-1, Mafb, Satb1, Nr2f1 (CoupTf1), Sp9, also appear to be downregulated in discrete 308 populations. This also includes a downregulation of Bcl11b (Ctip2) in both hippocampal 309 and cortical NGFCs, a gene recently linked to regulation of NGFC morphology and func-310 tion [65]. Among the few transcriptional regulators upregulated are Sirt2, Elmo1, Zcchc12, 311 none of which have been characterized in the context of MGE function (Figure $4B_{ii}$). Sirt2 312 is an established transcriptional repressor [66, 67] that may regulate the repression of sev-313 eral target genes in an MGE-specific manner, and *Elmo1* has been previously characterized 314 during the activity-dependent migration of CGE subtypes [21]. Finally, a recent study has 315 predicted that the expression of Zcchc12 correlates with slower intrinsic firing among hip-316 pocampal CA1 interneurons [43]. This suggests that increased Zcchc12 expression might 317 regulate the expression of synaptic genes enabling reduced intrinsic excitability in the MGE 318 subsets. Related to such putative decreased excitability in the MGE-derived interneurons, 319 among the activity-regulated TFs, we observe broad downregulation of Jun, Egr1, Fos, Fosb, 320 Arc, Satb1, Arnt2 across all classes of MGE in both brain regions (Figure4 C_{ii}). While 321 most of these are well-established activity-regulated TFs, Arnt2 has been recently described 322 to partner with Npas4, downstream of Ca^{2+} signaling in response to neuronal activity [68]. 323 Unsurprisingly, the Npas-family members Npas1/3/4 are also downregulated in discrete 324 MGE subtypes. 325

Impaired NMDAR signaling alters region-specific MGE subtype marker ex pression

Several GABAergic/MGE markers were mis-regulated upon Grin1-ablation (Figure5A). For example, genes S100a0, Pthlh, Hcrtr2 that are normally expressed in SST, PV and NGFCs respectively, are upregulated in the same clusters of MGE- $Grin1^{fl/fl}$ (Figure5B_i), indicating a misexpression in a subtype-specific manner. Next, while certain genes such as Reln, Tenm1 are broadly downregulated across MGE classes, some genes like Thsd7a show an upregulation in certain classes but a down regulation in

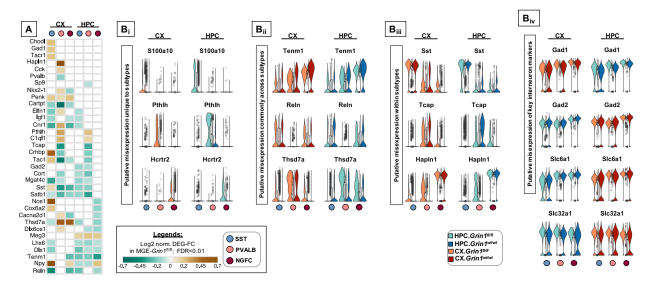


Figure 5: Fig. 5 | Differential expression of interneuron marker genes across subtypes upon *Grin1*-ablation A, Heatmap of log2 FC of significant DEGs in cortical and hippocampal MGE cardinal subtypes, showing a subset notable MGE marker genes. Representative box-violin plots of top differentially expressed genes from the above that represent B_i , a misexpression unique to a single MGE- $Grin1^{fl/fl}$ subtype; B_{ii} , a misexpression across all MGE- $Grin1^{fl/fl}$ subtypes; B_{iii} , a misexpression between MGE- $Grin1^{fl/fl}$ subtypes. B_{iv} , Representative box-violin plots of fundamental interneuron markers.

the other classes (Figure 5B_{ii}). Interestingly, a few genes that are normally abundant in 334 one MGE class, appear to be misexpressed in another MGE class where they are not abun-335 dant. For instance, Tcap, that is normally expressed in PV cells, in addition to being 336 decreased in PV cells, is upregulated in NGFCs in both cortex and hippocampus. Simi-337 larly, Hapln1 expression which is typically limited to NGFCs, is upregulated in PV subsets 338 (Figure 5B_{iii}). Lastly, we observed an upregulation in the *Gad1* and *Slc32a1* (vesicular 339 GABA transporter, vGAT) and a downregulation in *Gad2* and *Slc6a1* (Na⁺-Cl⁻dependent 340 GABA transporter, GAT1), corresponding with GABA synthesis and reuptake machineries 341 respectively (Figure $5B_{iv}$). Taken together, these data indicate that *Grin1*-ablation alters 342 region-specific MGE subtype numbers, and subtype marker expression indicative of altered 343 subtype identities. 344

NMDAR signaling regulates MGE subtype-specific expression of neurodevel opmental disorder risk genes

Interneuron-centric disease etiology is an emerging centrality in multiple psychiatric dis-347 orders [23]. Thus, we questioned whether the *Grin1* ablation induced DEGs presently iden-348 tified correlate with disease etiology. Disease-ontology based Ingenuity Pathway Analysis of 349 the DEGs showed significant over-representation of genes implicated in 'Schizophrenia', 'Psy-350 chiatric disorders' and 'Movement disorders', among other cellular impairments involving 351 aberrant morphology of neurons (Figure 6A, Supplemental Table 4). To independently 352 examine the DEGs for potential enrichment for neurodevelopmental disorders, we obtained 353 the risk genes for schizophrenia (Sz) and autism spectrum (As) from the SZDB [69] and 354 SFARI [70] databases respectively. These databases curate and rank disease-relevant gene 355

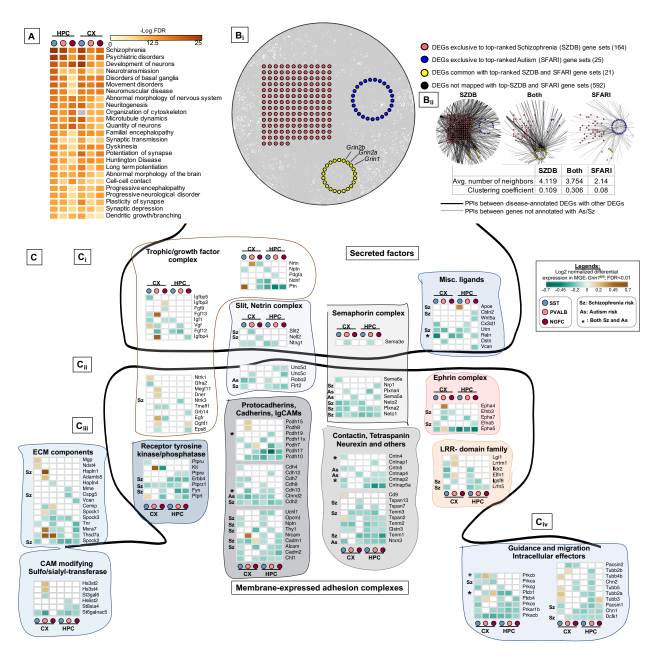


Figure 6: Aberrant *Grin1*-signaling result in misexpression of high-risk Sz genes A, Ingenuity Pathway Analysis of significantly overrepresented disease pathways in each MGE-subtype. \mathbf{B}_{i} , Global protein-protein interaction (PPI)map among all differentially expressed genes (DEGs). Red circles indicate the DEGs annotated to be top-ranked Sz-risk genes; Blue circles indicate the DEGs annotated to be top-ranked As-risk genes; Yellow circles indicate the DEGs annotated with both Sz and As-risk genes. Black circles in the periphery indicate the DEGs not annotated with high-risk Sz/As genes. The PPIs between DEGs indicated in grey lines. \mathbf{B}_{ii} , PPIs between Sz / As / dually enriched clusters, and other genes. The PPIs between disease-annotated DEGs and other disease-annotated DEGs or with other nonannotated DEGs are indicated in black lines. ThePPI between non-annotated DEGs indicated in grey lines. **C**, Heatmap of log2 FC of significant DEGs in cortical and hippocampal MGE cardinal subtypes, showing a subset of \mathbf{C}_{i} , secreted trophic factors and secreted ligands and guidance cues. \mathbf{C}_{ii} , membrane-bound synaptogenic receptors and cell adhesion molecules (CAMs) \mathbf{C}_{iii} , Extracellular Matrix (ECM) components and matrix modifying enzymes. \mathbf{C}_{iv} , Intracellular effectors of guidance and synaptogenic cues.

sets, based on multiple evidence sources including genome-wide association studies, copy-356 number variations, known human mutations and other integrative analyses. In particular, 357 we mapped the DEGs with the top-ranked genes from these disease datasets (see methods for 358 details). While 592 DEGs could not be mapped with either disease genes, 25 genes mapped 359 exclusively with the SFARI-AS gene list, 164 genes mapped exclusively with the SZDB-Sz 360 gene list and 21 genes mapped with both datasets (Figure6B_i, Supplemental Table5). 361 It is now well-established that several neurodevelopmental disorders exhibit a high degree of 362 converging molecular pathways employing proteins that exist in physical complexes [71-74]. 363 Therefore, we examined whether these disease-associated DEGs are known to form pro-364 tein complexes between each other, by mapping curated protein-protein interaction (PPI) 365 datasets for all 802 DEG products. Indeed, we observed that >95% of disease-annotated 366 DEG products are known to exist with PPIs, while only $\sim 75\%$ of DEG products not anno-367 tated with Sz/As are known to exist with PPIs (Supplemental Table5C). Interestingly, 368 despite not mapping directly with the high-ranked disease gene sets, the remaining 592 369 genes are observed to exist in tightly-knit PPIs with the disease annotated genes. How-370 ever, the PPIs mapped with SZDB form the most interconnected clusters in comparison to 371 the SFARI-mapped PPI network (Figure $6B_{ii}$), as indicated by relatively higher clustering 372 coefficient. This indicate that members of the DEGs here identified share physical, and 373 functional pathways in MGE-derived interneurons, contributing towards disease etiology. 374

Among the 210 DEGs mapped with to Sz and As, 45 genes are established regulators 375 of axon path-finding, synapse formation and assembly, while 38 members are established 376 regulators of membrane excitability and neuronal firing. Because both of these gene classes 377 are intimately associated with interneuron function, we examined these classes in detail. We 378 observed multiple classes of secreted ligands and cognate receptor families corresponding 379 to semaphorin, netrin, slit, chemokine and growth factors, and their intracellular effectors 380 that are downregulated upon MGE-Grin1-ablation (Figure $6C_{i,iv}$). These include Ntng1, 381 Sema3e, Slit2, Cx3cl1, and some of their receptors, Unc5c, Nrp1, Neto1/2, Robo2 that are 382 decreased in a MGE-class-specific manner. We observed Fqf13 that was recently demon-383 strated to mediate MGE-subtype specific synapse assembly [75], to be upregulated in cortical 384 PV cells, but downregulated in cortical SST, while Apoe to be upregulated in hippocam-385 pal SST cells. In addition to synaptic assembly molecules, we observed DE in a variety 386 of synaptic adhesion molecules, corresponding to protocadherin, cadherin, ephrin and con-387 tactin families (Figure $6C_{ii}$). Notably, we also observed a downregulation of *Erbb*4 across 388 all hippocampal MGE-subtypes. Lastly, we observed increased expression of extracellular 389 matrix components Mqp, Ndst4, Hapln1, Adamts5, Mxra7, Thsd7a and the matrix modify-390 ing enzymes Hs3st2/4 in cortical SST/PV subtypes (Figure6C_{iii}). 391

Among the regulators of neuronal excitability, we observed a downregulation of multiple members of postsynaptic glutamate receptor subunits, GABA receptors and their associated partners (Figure6-Supplement1B_{ii}). Interestingly, while we noted a broad downregulation of several members of potassium and sodium channel subunits, a few discrete members of the *Kcn*-families were upregulated in cortical PV and NGFC subtypes. Finally, we also observed multiple members of presynaptic GABA synthesis, release and uptake machineries including *Gad1*, *Syt2/10*, and *Slc6a1* differentially expressed in discrete MGE subtypes (Figure6, Supplement1 B_i). Collectively, these findings highlight the centrality of MGE-expressed *Grin1*-signaling during synapse formation and connectivity, which when aberrantly expressed, can lead to neurodevelopmental disorders.

402 Discussion

403 Centrality of MGE-derived interneuron-expressed NMDARs from juvenile 404 brain

NMDARs serve as critical activity dependent signaling hubs for myriad neuronal func-405 tions due to their innate ability to directly link network dynamics to cellular calcium events, 406 and associated transcriptional coupling. Such NMDAR-dependent excitation-transcription 407 coupling is widely established in glutamatergic neurons [76], and in specific interneurons 408 using candidate approaches [77] within mature circuits. However, the detailed unbiased 409 evaluation of the transcriptional landscape of NMDAR signaling within interneurons in de-410 veloping circuits undergoing refinement is lacking. Our study provides the first systematic 411 "fingerprinting" of the transcriptional coupling associated with NMDAR signaling, exclusive 412 to MGE-derived interneurons, providing a road map for examining NMDAR regulation of 413 MGE-derived interneurons in a subtype specific manner. 414

Our unbiased transcriptional profiling approach indicates that developmental NMDAR 415 signaling participates in MGE-derived interneuron specification by regulating the expres-416 sion of transcription factors (67 genes), synaptogenic (53 genes) and connectivity fac-417 tors/adhesion molecules (61 genes), and regulators of membrane excitability (78 genes), 418 among the 802 DEGs in interneurons (Figure 7). We employed bioinformatic analy-419 ses to examine whether system-wide downregulation of target genes can be attributed to 420 transcription-repression elements. Indeed, we identify a set of 10 transcriptional regulators 421 that commonly recognize the DNA-motifs present in the identified DEGs, including the 422 master-repressor *Rest.* Future studies are needed to examine the role of these putative re-423 pressor and repression motifs that have not been previously associated with MGE-specific 424 transcription. However, based on broad transcriptional downregulation of target genes, we 425 can make several predictions that should guide future investigations. 426

427 Shaping interneuron identity and granularity amongst subtypes

Interneuron development from MGE is replete with combinatorial expressions of numer-428 ous transcription factors, leading to diversity [78, 79]. Several transcription factors that are 429 impacted by Grin1-signaling are established regulators of MGE fate, subtype abundances 430 and identities (34 genes) including Nkx2-1, Lhx6, Dlx1, Dlx6, Maf, Mafb, Mef2c, Etv1, 431 Npas1, Npas3 and Sp9 [34, 56, 80–83] [84]. While the scRNAseq landscapes of interneurons 432 predict several transcriptomic features that would classify them as distinct 'cell-types' or 433 cell-states' [49, 85, 86] [50] the precise mechanisms responsible for such granularity is still 434 emerging [2]. It is possible that NMDAR signaling in the developing interneuron progenitors 435 may provide a combinatorial cue that will couple with innate genetic programs to gener-436 ate the diversity in interneuron subtypes. Indicating that the *Grin1*-lacking MGE-derived 437 interneurons have impaired subtype identities, we observe differential recoveries of the sub-438 types within SST, PV and NGFC in the scRNAseq assay (Figure2-Supplement2A,B,C). 430

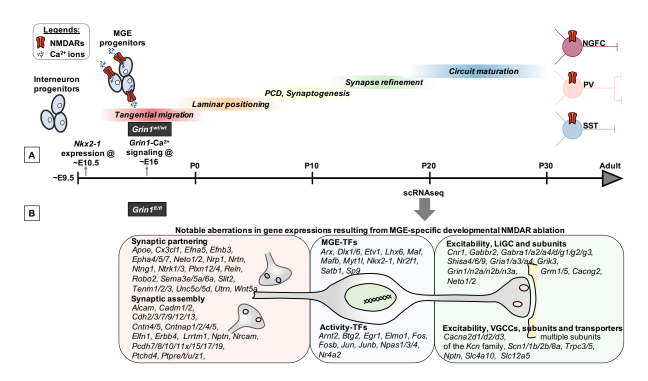


Figure 7: Transcriptional control of MGE development, synaptic partnering and excitability are mediated by NMDAR signaling A, Nkx2.1 expression appears at ~ED10.5, driving MGE subtype fate in interneuronal progenitors [42] [56]. Subsequently, their sequential developmental milestones towards circuit refinement appears to be under a combination of innate genetic mechanisms and neuronal activity. While the earliest Grin1 expression is reported at ~ED14 in developing brain [58] [57], MGE-specific Grin1mediated Ca^{2+} is recorded at ~ED16 [15]. However, the broad role played by interneuron-expressed NMDAR signaling during interneuron development until now is not well delineated. **B**, By driving Grin1-ablation using Nkx2.1-driven Cre-recombinase, we report the earliest developmental loss of NMDAR signaling, across MGE-derived interneuron subtypes. In particular, by performing scRNAseq assay in MGE-derived interneurons from the cortex and hippocampus of the mouse brain, we report a broad transcriptional aberration subsequent to loss of NMDAR-signaling. Notably, this expression abnormality involves numerous transcriptional factors, synaptogenic and regulators of interneuron excitability, that collectively estabish MGE subtype identities.

ED, Embryonic day; PD, Postnatal day; PCD, Programmed cell death; LiGC, Ligand-gated channel; VGCC, Voltage-gated calcium channel

Particularly, we find in the *Grin1*-ablated MGE-derived interneurons, the presence of diver-440 sified PVALB.3 populations that express marker genes such as *Prox1*, otherwise not robust 441 in our scRNAseq screen from the MGE- $Grin1^{wt}$. Additionally, by independent immunos-442 taining experiments, we observe a modest increase in cortical SST cell numbers along with 443 a modest decrease in cortical PV cells, in a manner similar to Maf: Mafb-mutants recently 444 reported [82]. However, detailed future studies are necessary to uncover whether/how the 445 NMDAR-dependent combinatorial transcriptional code works with innate mechanisms to 446 generate the diversity within PV/SST/NGFC subclasses. It would be intriguing to examine 447 whether these MGE-Grin1-null mice exhibit aberrations in the expression of these master 448 MGE-regulators such as Nkx2-1 and Lhx6 earlier in development. **44**C

450 Shaping interneuron subtype-specific synaptic assembly and connectivity

What is the biological context of differential expression of the TFs, in the juvenile fore-451 brain when MGE-derived interneuron fate is assumed to be already sealed, and subtype 452 identities established? It is emerging that some of these TFs are continually required for the 453 maintenance of MGE fate, post development [87]. One of the ways the TFs maintain MGE 454 subtype fate into adulthood, is by controlling the expression of genes that are essential for 455 ongoing interneuron function. Accordingly, we predict that NMDAR-dependent expression 456 of synaptogenic and synaptic partnering molecules regulate the assembly of synapses with 457 appropriate targets. Secreted semaphorin, ephrin, slit, netrin and neurotrophin-based sig-458 naling systems have been investigated in GABAergic neurons, during axonal pathfinding, 459 and cell migration [88–95]. However, only recently have inroads been made into delineat-460 ing their expression, and interaction with appropriate receptor systems in target synapses 461 during accurate synaptogenesis. In addition, the NMDAR-dependent expression of synap-462 tic adhesion molecules will further promote stability of newly formed synapses. Here, the 463 mis-expression of diverse secreted cues, their receptors and adhesion molecules by MGE sub-464 types during *Grin1*-ablation, provides unique insight into the molecular diversity employed 465 during synapse establishment. Our findings also reveal numerous candidates for examining 466 subtype specific synapse assembly, which are centrally regulated by NMDAR signaling. Of 467 particular interest are the family of protocadherins that are reported recently to be com-468 monly downregulated in cortical interneurons generated from Sz patient-derived induced 469 pluripotent stem cells [96]. 470

Subsequent to synapse formation nascent connections remain susceptible to strength 471 modifications according to neuronal activity. Again, NMDAR-signaling in MGE-derived in-472 terneurons seems to regulate this process by the transcriptional regulation of the expressions 473 of both presynaptic and postsynaptic members, including excitatory and inhibitory synap-474 tic molecules and their auxiliary subunits, as well as presynaptic GABA release machinery 475 molecules such as Cplx1/2, Stx1b, Rab3c. However, most dramatic is the massive down reg-476 ulation of several members of the potassium channel subunits and their auxiliary subunits 477 across MGE subtypes, with the exception of an upregulation of a few Kcn-genes in cortical 478 PVs and NGFCs. While the precise impact of the diverse changes in these genes on MGE 479 firing are currently unclear, the pattern of expression of the activity-dependent transcription 480 factors provides us an indication. 481

Notable activity-dependent TFs such as Jun, Eqr1 are downregulated across all MGE 482 subtypes, while Fosb, Fos, Arx are down regulated across all hippocampal MGEs, and 483 Satb1, Arnt2 are downregulated across all cortical MGEs. In addition, Npas4, an established 484 early-response TF [97–99] activated upon neuronal activity and Ca²⁺ influx in MGE-derived 485 interneurons [100], was downregulated in cortical NGFCs upon Grin1-ablation. Etv1 was 486 previously demonstrated to be an activity-dependent TF that inversely correlates Ca^{2+} in-487 flux, regulating the identity of a subset of PV-interneurons [80]. Remarkably, we observe an 488 increase in *Etv1* expression in cortical PV cells. Lastly, *Ostn* was recently established as an 489 activity-regulated secreted factor [101], and we observed Ostn to be downregulated specifi-490 cally in cortical PV subtypes (Figure 4C $_{ii}$). Together, these changes are consistent with 491 reduced neuronal activity in MGE subtypes upon *Grin1*-ablation, consistent with previous 492

reports indicating that NMDAR-antagonists can directly reduce the activity of GABAergic interneurons in adult mice [27]. Interpreting the differential expressions of activitydependent genes during scRNAseq has been challenging, particularly, when these genes could get activated by the very process involved in cell dissociation and sorting [102, 103]. However, our use of activity-blockers and actinomycin-D throughout our MGE- $Grin1^{wt}$ and MGE- $Grin1^{fl/fl}$ scRNAseq pipelines [46], gives confidence that the differential expressions of activity-dependent TFs reflect biological relevance.

500 NMDAR signaling in NGFCs

Among the MGE subtypes, the PV and SST interneurons are traditionally widely stud-501 ied in comparison to the dendrite-targeting NGFC subtypes (that include the Ivy cells). In 502 the present study we provide the first detailed molecular insight into the cortical and hip-503 pocampal NGFCs, subsequent to NMDAR ablation. We anticipated that these cell types 504 could be particularly susceptible to loss of NMDARs, since we previously reported that 505 NGFCs exhibit the most robust synaptic NMDAR conductances among the MGE sub-506 types [12]. Intriguingly, while the cortical NGFCs had comparable numbers of both total 507 and unique DEGs with respect to other cortical MGE-derived interneurons (Figure 3B), we 508 observed far fewer total and NGFC-specific DEGs in the hippocampus, compared to other 509 hippocampal MGEs. However, based on the scRNAseq cell type recoveries, we predict an 510 elaboration of NGFC.1, and a reduction in the NGFC.2 subtype upon *Grin1*-ablation. Fi-511 nally, NGFCs exhibited dendritic arborization impairments subsequent to impaired NMDAR 512 signaling [9, 11]. Indeed, we observe 49 genes among the DEGs (Supplemental Table1 513 that have established roles in regulating neuronal cytoskeleton and associated signaling,) 514 likely mediating the observed dendritic impairments in NGFCs. 515

⁵¹⁶ Developmental NMDAR ablation in interneurons and schizophrenia

Impaired NMDAR function observed during human NMDAR gene mutations [104], and 517 anti-NMDAR-encephalitis [105] results in a wide range of neuropsychiatric disorders in-518 cluding autism spectrum disorders [106, 107], intellectual disability [108], psychosis [109], 519 epilepsy and associated comorbidities [110, 111]. While broadly aberrant NMDAR signaling 520 in neurons is thought to underlie a wide range of these neurological disorders, an interneuron-521 centric developmental NMDAR aberration is emerging central to schizophrenia-related syn-522 dromes. Indeed, in the present study, disease mapping of the DEGs using high-ranked 523 SZDB-Sz and SFARI-As datasets indicate that many more DEGs map with the Sz than 524 the As database. Moreover, these disease-relevent DEGs exist in physical and functional 525 complexes with other DEGs that are not directly mapped to the Sz database. We used only 526 stringent, high-ranked disease genes from the database that pass several disease-relevant cri-527 teria. However, there are other DEGs that still map to lower-ranked Sz and As datasets that 528 are 'non-annotated' in present study. While our study can be argued as an 'extreme' case 529 of NMDAR hypofunction in MGE-derived interneurons, it provides a starting point high-530 lighting the centrality and broad range of interneuronal NMDAR-transcriptional pathways 531 during development. 532

A multitude of studies implicate NMDAR-hypofunction specific to PV cell types as a central underlying feature of schizophrenia etiology [112, 113]. However, the measurable NM-DAR conductances within PV interneurons are relatively small in comparison to other MGE

subtypes [12]. Additionally, NMDA signaling in non-PV interneuron subtypes drives robust
dendritic inhibition in pyramidal neurons [114, 115]. Moreover, while NMDAR-ablation in *Pvalb*-Cre lines produces other behavioral deficits unrelated to the Sz-like phenotypes [30, 33]
, a developmental, but not adult-onset *Grin1*-ablation in *Ppp1r2*-Cre line [29] that targets
a subset of PV interneurons among other subtypes [116], recapitulates core Sz-like phenotypes. Lastly, studies that map interneuron subtypes to Sz-like phenotypes indeed support
the role of different interneuron classes beyond PV cells towards disease etiology [28, 33].

Integrating these ideas and based on findings from the present study, we propose the 543 following: (i) Despite a smaller NMDAR conductance in PV interneurons, we observe a ro-544 bust transcriptional coupling via NMDARs, as observed by several distinct gene expression 545 abnormalities in this cell type relevant to human Sz. Therefore, PV-expressed NMDARs pri-546 marily serve to regulate transcriptional coupling, mediating the abundances of PV-subtype 547 abundances. (ii) The developmental window for NMDAR loss of function is particularly 548 important because, its transcriptional regulation maintains the correct synaptogenic and 549 assembly cues, which when lost, lead to disease causing-impaired connectivity. Perhaps, 550 in the $Grin1^{fl/fl}$: Pvalb-Cre mouse line, the Grin1-ablation occurs only at a developmen-551 tal window when synaptic connectivity is sufficiently complete, explaining why the animal 552 model does not lead to profound Sz-like impairments. (iii) The dendrite targeting SST and 553 NGFC interneurons also exhibit robust NMDAR signaling and transcriptional coupling. 554 During aberrant NMDAR-transcriptional coupling, it is therefore likely that impaired den-555 dritic connectivity and inhibition onto pyramidal neurons also contributes towards disease 556 etiology. Therefore, our dataset provides credence to interneuronal subtype-specific granu-557 larily, connectivity and excitability, all playing combinatorial and mutually-supporting roles 558 during disease etiology. 559

Taken together, our study presents a rich resource, laying the road map for systematic examination of NMDAR signaling in interneuron subtypes, by providing multiple molecular targets for examination in both normal and impaired circuits.

563 Materials and methods

564 Contact for Reagent and Resource Sharing

Further information and requests for resources and reagents should be directed to and reasonable requests will be fulfilled by the Lead Contact, Chris McBain (mcbainc@mail.nih.gov).

568 Animals

All experiments were conducted in accordance with animal protocols approved by the Na-560 tional Institutes of Health. The Nkx2.1-Cre driver line (C57BL/6J-Tg(Nkx2-1-Cre)2Sand/J; 570 Cat. No. 008661 | Nkx2.1-Cre, Floxed Grin1 mouse line (B6.129S4-Grin1^{tm2Stl}/J; Cat. No. 571 005246 | fNR1) and Ai14 reporter mouse (B6.Cg-Gt (ROSA)26Sor^{tm14(CAG-tdTomato)Hze/J}; 572 Cat. No. 007914 | Ai14, Ai14D or Ai14(RCL-tdT)-D), purchased from the Jackson Labora-573 tory, were used to generate the MGE-derived interneuron-specific $Grin1^{fl/fl}$ line. Littermate 574 MGE-Grin $1^{wt/wt}$ controls, and both male and female mice were used during this study. Mice 575 were housed and bred in conventional vivarium with standard laboratory chow and water in 576

standard animal cages under a 12hr circadian cycle. Genotyping of the mice were performed
as indicated in the appropriate Jackson Laboratory mice catalog.

579 Single-cell dissociation and FACS

P18-20 juvenile Nkx2-1-Cre: Grin1^{wt/wt}: TdT⁺ and Nkx2-1-Cre: Grin1^{fl/fl}: TdT⁺ mice 580 were used for single-cell sequencing experiments. All mice were anesthetized with isoflu-581 rane and then decapitated. Brain dissection, slicing and FACS sorting were carried out 582 as described [43, 44], with slight modifications. NMDG-HEPES-based solution was used 583 in all steps to enable better recovery of the cells [45] during FACS sorting and single-cell 584 bar coding. Briefly, the brain sectioning solution contained NMDG-HEPES-based high-585 Mg^{2+} cutting solution contained 93 mM NMDG, 2.5 mM KCl, 1.2 mM NaH₂PO₄, 30 mM 586 NaHCO₃, 20 mM HEPES, 25 mM glucose, 5 mM sodium ascorbate, 3 mM sodium pyruvate, 587 2mM Thiourea, 10 mM MgSO₄*7H₂O, and 0.5 mM CaCl₂*2H₂O; it was adjusted to pH 7.4 588 with 12.1N HCl, an osmolarity of 300-310 mOsm, and carbogenated (mix of 95% O₂ and 589 5% CO₂) before use. This solution was chilled and the process of sectioning were conducted 590 on a ice-chamber in the vibratome. 591

3-4, Nkx2-1-Cre: Grin1^{wt/wt}: TdT⁺ or Nkx2-1-Cre: Grin1^{fl/fl}: TdT⁺ mice were pro-592 cessed on consecutive days for single-cell sequencing experiments. TdT negative animals 593 were processed in parallel for initially setting FACS gate for the Tomato-channel. Across 594 the replicates, 10XMGE-Grin1-WT and 6XMGE-Grin1-null animals were used for the scR-595 NAseq. Coronal slices containing frontal cortex and hippocampus (350mM) were cut using 596 VT-1000S vibratome (Leica Microsystems) in cold NMDG-HEPES-based high-Mg²⁺ cut-597 ting solution. Slices were recovered in the same solution at 20°C for 30 minutes during 598 when, they were visually inspected under fluorescence microscope and micro dissected, all 599 under constant carbogenation. The recovery and microdissection were conducted in the 600 NMDG-HEPES high-Mg²⁺ solution supplemented with 0.5μ M tetrodotoxin (TTX), 50 μ M 601 DL -2-Amino-5-phosphonopentanoic acid (APV) and 10 μ M Actinomycin-D (Act-D). 602

Cell dissociation was performed using the Worthington Papain Dissociation System 603 (LK003150) according to manufacturer instructions with minor modifications. Briefly, 604 single-cell suspensions of the micro dissected frontal cortices or hippocampus were prepared 605 using sufficiently carbogenated dissociation solution (containing Papain, DNAse in Earle's 606 Balanced Salt Solution, EBSS), supplemented with $1\mu M$ TTX, 100 μM APV and 20 μM 607 Act-D. After a 60 min enzymatic digestion at 37°C, followed by gentle manual trituration 608 with fire-polished Pasteur pipettes, the cell dissociates were centrifuged at 300g for 5 min-609 utes at 20°C, and the supernatants were discarded. The enzymatic digestion was quenched 610 in the next step by the addition of ovomucoid protease inhibitor. Albumin density gradi-611 ent was performed on the pellets, using a sufficiently carbogenated debris removal solution 612 (containing albumin-ovomucoid inhibitor, DNAse in EBSS). The resulting cell pellets were 613 resuspended in 1ml FACS buffer containing 10% FBS, $10U/\mu$ l of DNAse, 1 μ M TTX, 100 614 μ M APV and 20 μ M Act-D in a 50:50 mix of carbogenated EBSS: NMDG-HEPES-based 615 cutting saline (with $1 \text{mM MgSO}_4*7\text{H}_2\text{O}$, it is important to not use High-Mg²⁺ in the FACS 616 buffer, as it interferes with the subsequent 10X scRNAseq reaction). Cells were placed in 617 polystyrene tubes (Falcon 352235) on ice during the FACS. 618

⁶¹⁹ For single cell sorting of TdT⁺ expressing cells by FACS, resuspended cell dissociates

were filtered through 35mm cell strainer (Falcon 352235) to remove cell clumps. The single 620 cell suspensions were then incubated with 1mg/ml DAPI (1:500, Thermo Scientific 62248) 621 and 1mM DRAQ5 (Thermo Scientific 62251) at 4°C for 5 minutes to label dead cells and live 622 cells respectively. Samples were analyzed for TdTomato expression and sorted using a MoFlo 623 Astrios EQ high speed cell sorter (Beckman Coulter). TdT-negative cells were used as a con-624 trol to set the thresholding FACS gate for the detection and sorting of the Ai14-TdTomato-625 expressing cells, and the same gate was then applied for all subsequent experiments. Flow 626 data analysis and setting of sorting gates on live (DAPI-negative, DRAQ5-positive) and 627 Ai14-TdTomato-expressing cells were carried out using Summit software V6.3.016900 (Beck-628 man Coulter). Per sample/session, 20,000 – 40,000 individual cells were sorted into a FBS-629 precoated, Eppendorf LoBind Microcentrifuge tubes containing carbogenated 10ml FACS 630 buffer, that served as the starting material for 10X Genomics bar-coding. 631

632 10X Genomics Chromium

The cells were inspected for viability, counted, and loaded on the 10X Genomics 633 Chromium system, aiming to recover ~5000 cells per condition. 12 PCR cycles were con-634 ducted for cDNA amplification, and the subsequent library preparation and sequencing were 635 carried out in accordance with the manufacturer recommendation (ChromiumTM Single Cell 636 3' Library & Gel Bead Kit v2 and v3, 16 reactions). Sequencing of the libraries were 637 performed on the Illumina HiSeq2500 at the NICHD, Molecular Genomics Core facility. 638 Replicate 1 of the scRNAseq were performed using 10X v2 reaction from which, the cell 639 estimates, mean reads per cell (raw), median genes per cell respectively, are as follows Cor-640 tical WT: 1277, 149K, 4615; Cortical NULL: 181, 159K, 4826; Hippocampal WT: 2221, 92K, 641 2578; Hippocampal NULL: 404, 154K, 4903. Replicate 2 of the scRNAseq were performed 642 using 10X v3 reaction from which, the cell estimates, mean reads per cell (raw), median 643 genes per cell respectively, are as follows Cortical WT: 3851, 22.8K, 1536; Cortical NULL: 644 2898, 23.5K, 2759; Hippocampal WT: 4600, 23.6K, 850; Hippocampal NULL: 4436, 25.8K, 645 3143. Replicate 3 of the scRNAseq were performed using 10X v3 reaction from which, cell 646 estimates, mean reads per cell (raw), median genes per cell respectively, are as follows Cor-647 tical WT: 3960, 24.8K, 2870; Hippocampal WT: 3159, 26.9K, 2956. Representative quality 648 metrics from Replicate 2 are indicated in Figure 1-Supplement 1B, C, D, E. Demultiplexed 649 samples were aligned to the mouse reference genome (mm10). The end definitions of genes 650 were extended 4k bp downstream (or halfway to the next feature if closer), and converted 651 to mRNA counts using the Cell Ranger Version 2.1.1, provided by the manufacturer. 652

⁶⁵³ Data processing, analyses, visualization and differential expression testing

Processing (load, align, merge, cluster, differential expression testing) and visualiza-654 tion of the scRNAseq datasets were performed with the R statistical programming environ-655 ment [117] (v3.5.1) and Seurat package (v3.1.5, a development version of Seurat v3.1.5.9000 656 was used to generate violin plots in 2C and 5B) [47, 48]. Data set preprocessing, compari-657 son of WT- and NULL-Ai14 cells, canonical correlation analyses, and differential expression 658 of genes $(p_{adj} < 0.01)$ within the same cluster between WT- and NULL-Ai14 cells were 659 performed according to default Seurat parameters, unless otherwise mentioned. Quality 660 control filtering was performed by only including cells that had between 200-6000 unique 661 genes, and that had <30% of reads from mitochondrial genes. While the WT replicates 662

had no cells above 30% mitochondrial genes, only NULL replicates from both brain regions 663 exhibited 7-12% of cells above this threshold. Suggestive of inherent biological impact of 664 Grin1-ablation, we repeated the clustering and subsequent analyses without excluding any 665 cells. These analyses did not alter the clustering or skew the gene list. Clustering was per-666 formed on the top 25 PCs using the function FindClusters() by applying the shared nearest 667 neighbor modularity optimization with varying clustering resolution. A cluster resolution 668 of 1.0 was determined to be biologically meaningful, that yielded all known MGE cardinal 669 classes. Initial analyses were performed on the WT datasets separately (WT.alone), and sim-670 ilar set of analysis parameters were applied when the WT and NULL samples were merged 671 (WT.NULL.integrated) for subsequent differential expression testing. Phylogenetic tree re-672 lating the 'average' cell from each identity class based on a distance matrix constructed in 673 gene expression space using the BuildClusterTree() function. Overall, we identified 27, and 674 33 clusters using this approach in the WT.alone, and WT.NULL.integrated assays respec-675 tively. The WT.alone correspond to 11 MGE.GAD1/2 clusters (Figure1&2), while the 676 WT.NULL.integrated assay correspond to 12 clusters (Figure 5-Supplement.1). We first 677 searched for the top differential markers for each MGE subcluster using the FindAllMarkers() 678 function. The genes thus identified for the integrated data is presented in *Supplemental* 679 **Table1b.** Determination of MGE and non-MGE identities are performed based on existing 680 interneuron literature and other scRNAseq datasets [1, 43, 50, 53, 85, 87, 118–120]. The 681 labels from Figures 1 and 2 are matched with the top gene markers identified by the Find-682 AllMarkers() function and the similarly named clusters in Figures 1 and 2 have the same 683 identities. Lastly, for the integrated analyses and differential expression testing, we first 684 merged the identities of the subclusters SST.1-SST.6 and TH.1, and relabelled as SST sub-685 set; PVALB.1-3 relabelled as PVALB subset; and NGFC.1-2 relabelled as the NGFC subset 686 during subsequent analysis (Figure 3). 687

Differential gene expression testing were performed using the MAST package within the 688 FindMarkers function to identify the differentially expressed genes between two subclusters. 689 MAST utilizes a hurdle model with normalized UMI as covariate to generate the differential 690 fold changes [121], and is known to result in underestimation of the magnitude of fold 691 change (FC) [122]. Therefore, while applying a stringent false-discovery rate <0.01, we 692 determined the minimum FC based on the control gene Grin1, which is the target gene 693 knocked out in MGE-derived interneuron celltypes. Notably for *Grin1*, we had previously 694 demonstrated that the NGFCs which carry maximum NMDAR component among MGEs, 695 are devoid of NMDAR current at this comparable age [9]. In the present scRNAseq assay, 696 we observe a logFC for Grin1 ranging between -0.1 to -0.35 across both brain regions and 697 all MGE subtypes. Therefore, we determined a minimum logFC in our DEGs as ± 0.1 to be 698 meaningful. Previous studies have demonstrated the MAST approach for DEG testing to 699 be powerful in determining subtle changes in highly transcribed genes, and among abundant 700 populations, additional to under representing changes among weakly transcribed genes [121, 701 122]. Volcano plots and Heat maps for the DEG were generated using EnhancedVolcano 702 package [123] and Morpheus package https://software.broadinstitute.org/morpheus within 703 the R framework. 704

⁷⁰⁵ Pathway analyses, PPI network mapping and disease mapping

Ingenuity Pathway Analyses were conducted on the differentially expressed genes to gen-706 erate the molecular functional annotation and to identify the biological pathways and dis-707 ease pathways overrepresented. This tool was also used to annotate genes with their known 708 cellular functional classes. Additional Gene Ontology mapping and KEGG analyses were 709 conducted using ShinyGO [124]. Protein-protein interaction (PPI) mapping datasets from a 710 variety of curated databases [125–127] were conducted as previously described [72] [128] us-711 ing the Cytoscape [129] platform (v3.8.0). Schizophrenia risk genes integrated from various 712 sources including genome-wide association studies (GWAS), copy number variation (CNV), 713 association and linkage studies, post-mortem human brain gene expression, expression quan-714 titative trait loci (eQTL) and encyclopedia of DNA elements (ENCODE), were downloaded 715 from http://www.szdb.org/ [69]. Autism Spectrum Disorder risk genes integrated from var-716 ious sources were downloaded from Simons Foundation https://gene.sfari.org/ [70]. SZDB 717 genes that had a integrated total score of 3-6 (1419 genes, 22% out of 6387) were consid-718 ered 'high-risk' for DEG mapping (Supplemental Table5a). SFARI genes scored 1-2 with 719 accounting for a high strength of evidence (392 genes, 42% out of 943), were considered 720 'high-risk' for DEG mapping (Supplemental Table5b). Transcriptional factor motif en-721 richment search using the iRegulon [62] was also conducted using Cytoscape using default 722 parameters. 723

724 Immunostaining

All solutions were freshly prepared and filtered using 0.22μ m syringe filters for parallel 725 treatments of wildtype and MGE-Grin1-null groups. Adult mice of postnatal day (PD) 726 30/60/210 were Mice were deeply anesthetized with isoflurane and perfused transcardially 727 with 1X phosphate buffer saline (PBS) and followed by the fixative 4% paraformaldehyde. 728 The brains were post-fixed in the same fixative for overnight at 4 °C for the immunostaining 729 assays. Postfixed brains were serially dehydrated using 10%/20%/30% sucrose solutions at 730 4 °C. Coronal sections (50 μ m) were cut on a freezing microtome. Immunostaining was 731 performed on free-floating sections. Tissue sections were permeabilized and blocked in 1 \times 732 PBS + 1% bovine serum albumin + 10% normal goat serum + 0.5% Triton X-100 (Carrier 733 PB) at room temperature for 2 h, followed by incubation in primary antibodies, listed below, 734 diluted with $1 \times PBS + 1\%$ bovine serum albumin + 1% normal goat serum + 0.1% Triton 735 X-100 overnight at 4 °C. Tissue sections were then incubated with secondary antibodies, 736 listed below, diluted in Carrier Solution (1:1000), and DAPI (1:2000) at room temperature 737 for 1–2 h and mounted on Superfrost glass slides, and coverslipped using Mowiol mounting 738 medium and 1.5 mm cover glasses. 739

740 Antibodies

The following primary antibodies were used: mouse anti-PV (1:1000; Sigma-Aldrich
Cat# P3088, RRID: AB_477329), rat anti-SST (1:1000; Millipore Cat# MAB354, RRID:
AB_2255365). Secondary antibodies were conjugated with Alexa Fluor dyes 488 or 633
(1:1000; Thermo Fisher Scientific).

⁷⁴⁵ Image acquisition and analysis

Mouse brains from 4–8 different animals were used for each condition, and section depth were matched between the genotypes for parallel immunostaining. Fluorescent images were captured using the 10X objective of a Nikon Widefield Fluorescence, Spinning Disk Confocal

microscope. For all slices with immunostained or genetically reported signal, 50 μ m thin 749 sections were imaged using 10x/0.45 CFI PlanApo objective (imaging settings: Numerical 750 Aperture 0.75, bit depth 16-bit, Exposure 100ms). Confocal stacks were stitched using NIS 751 Elements (Nikon) before importing them into Imaris software (Bitplane, version 9.2). Cell 752 bodies were marked in Imaris software using the 'Spots' function. Nkx2-1-Cre:TdT⁺ RFP+, 753 PV+ cell bodies were detected using the automatic function, with a signal detection radius 754 of 10 μ m. The Imaris 'Quality' filter was set above an empirically determined threshold to 755 maximize the number of detected cells while minimizing observed false positives. SST+ cell 756 bodies were marked manually using the Imaris 'Spots' function. ROI 3D borders around 757 hippocampus or cortex, drawn manually using the Imaris function 'Surfaces'. Spots were 758 then split within each ROI using the Imaris function 'Split Spots'. Overlap of RFP+ cells 759 with other markers (PV, SST) was addressed by filtering the RFP+ Spots above an em-760 pirically determined threshold intensity in the channel relative to the marker of interest. 761 Each image with an automatic analysis by Imaris was checked by an expert and incorrectly 762 identified cell bodies where refined if required. In Figure 5A, B Error bars reflect standard 763 error of mean; Two-tailed unpaired t-test was performed using Prism8. 764

765 Author contributions

VM and CJM conceived the project. VM, TJP and CJM designed the experiments, DM performed FACS sorting and analysis. VM, YZ, TJP performed 10X scRNAseq. VM, AM, CJR, CE, RD performed 10X scRNAseq bioinformatic analyses. VM and AP conducted immunuflourescent staining, imaging and analysis. CJM supervised the study. VM and CJM wrote the paper and all authors edited the manuscript.

771 Acknowledgment

This work was supported by *Eunice Kennedy Shriver* NICHD Intramural Award to CJM. We thank Steven L. Coon, Tianwei Li and James R. Iben at the Molecular Genomics Core, NICHD, for RNA sequencing and bioinformatics support. We thank Vincent Schram and Lynne Holtzclaw of the NICHD Microscopy and Imaging Core for imaging support, and we thank Carolina Bengtsson Gonzales for assistance with improving cell viability during dissociation and FACS. We also thank Xiaoqing Yuan, Steven Hunt, Daniel Abebe for assistance with animal colony maintenance.

779 References

- [1] K. A. Pelkey, R. Chittajallu, M. T. Craig, L. Tricoire, J. C. Wester, C. J. McBain, Hippocampal GABAergic Inhibitory Interneurons, Physiological Reviews 97 (4) (2017) 1619–1747. doi:10.1152/physrev.00007.2017.
- 783 URL https://dx.doi.org/10.1152/physrev.00007.2017
- [2] B. Wamsley, G. Fishell, Genetic and activity-dependent mechanisms underlying interneuron diversity,
- 785 Nature Reviews Neuroscience 18 (5) (2017) 299–309. doi:10.1038/nrn.2017.30.
- 786 URL https://dx.doi.org/10.1038/nrn.2017.30

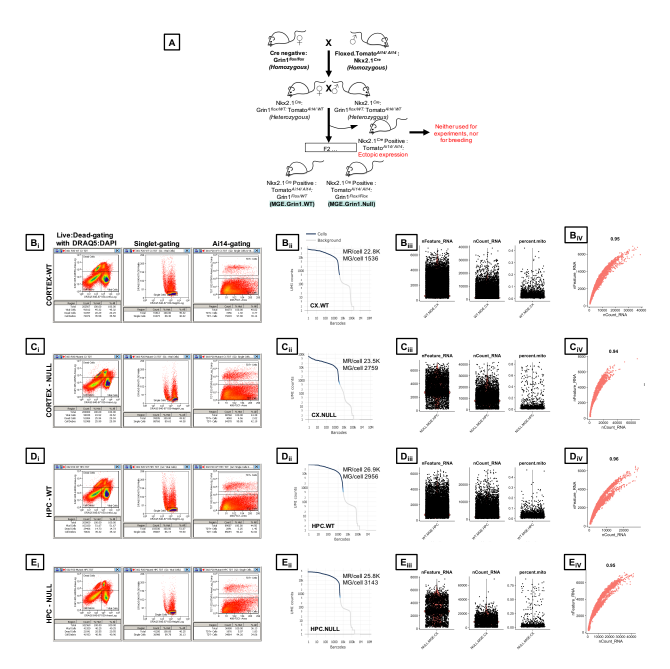
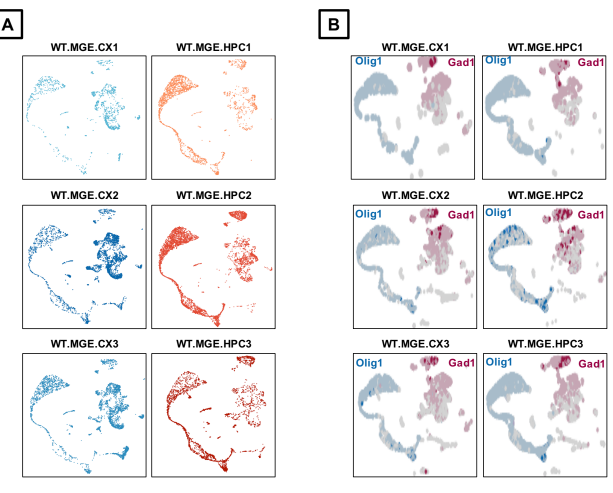


Figure 8: (Figure1, Supp.1) | Schematic overview and quality control for scRNAseq A, Breeding strategy, $\mathbf{B_i}, \mathbf{C_i}, \mathbf{D_i}, \mathbf{E_i}$, Representative FACS gates to sequentially isolate live: dead cells using DAPI: DRAQ5 staining, singlet-gating and TdT⁺-reporter gating to obtain reporter-positive MGE-derived interneurons from frontal cortex and hippocampus. $\mathbf{B}_{ii}, \mathbf{C_{ii}}, \mathbf{D}_{ii}, \mathbf{E}_{ii}$, Barcode Rank Plots for cells from WT and NULL mice, demonstrating separation of cell-associated barcodes and those associated with empty partitions. UMI, unique molecular identifier; MR, Mean Reads; MG, Median Genes. $\mathbf{B}_{iii}, \mathbf{C}_{iii}, \mathbf{D}_{iii}, \mathbf{E}_{iii}$, Distributions of the total number of genes, percentage of mitochondrial genes and UMIs per cell in control mice $\mathbf{B_{iv}}, \mathbf{C}_{iv}, \mathbf{D}_{iv}, \mathbf{E}_{iv}$, Pearson correlation coefficient of the distributions of the total number of genes and the UMI



Biological replicates

Biological replicates

Figure 9: (Figure 1, Supp.2) | Biological replicates Representative UMAP plots of the 3 biological replicates from cortex and hippocampus indicates \mathbf{A} , similar clustering, and \mathbf{B} , similar expression profiles of *Nkx2-1*-derived, *Gad1*-expressing MGE-derived interneurons and *Nkx2-1* derived, *Olig1*-expressing oligodendrocytes.

- [3] L. Tricoire, K. A. Pelkey, B. E. Erkkila, B. W. Jeffries, X. Yuan, C. J. McBain, A Blueprint for the Spatiotemporal Origins of Mouse Hippocampal Interneuron Diversity, Journal of Neuroscience 31 (30) (2011) 10948–10970. doi:10.1523/jneurosci.0323-11.2011.
- 790 URL https://dx.doi.org/10.1523/jneurosci.0323-11.2011
- [4] L. Tricoire, K. A. Pelkey, M. I. Daw, V. H. Sousa, G. Miyoshi, B. Jeffries, B. Cauli, G. Fishell, C. J. McBain, Common Origins of Hippocampal Ivy and Nitric Oxide Synthase Expressing Neurogliaform Cells, Journal of Neuroscience 30 (6) (2010) 2165–2176. doi:10.1523/jneurosci.5123-09.2010.
- 794 URL https://dx.doi.org/10.1523/jneurosci.5123-09.2010
- [5] F. M. Krienen, M. Goldman, Q. Zhang, R. D. Rosario, M. Florio, R. Machold, A. Saunders,
 K. Levandowski, H. Zaniewski, B. Schuman (2019).
- [6] L. Overstreet-Wadiche, C. J. McBain, Neurogliaform cells in cortical circuits, Nature Reviews Neuroscience 16 (8) (2015) 458-468. doi:10.1038/nrn3969.

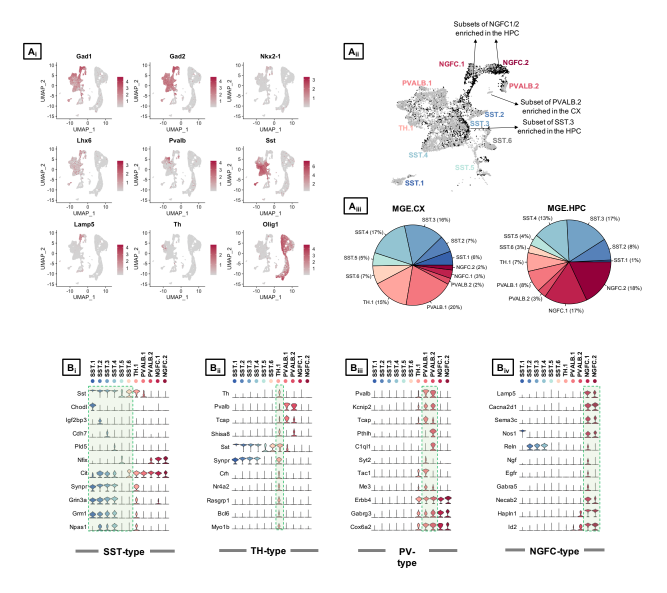


Figure 10: (Figure 1, Supp.3) | Select marker gene expression across the subtypes of merged cortical and hippocampal MGE-*Grin1* $^{wt/wt}$ \mathbf{A}_i , UMAP representation of cardinal MGE markers genes in the cortical and hippocampal merged dataset. \mathbf{A}_{ii} , UMAP representation colored by region, highlighting the region-specific enrichments of MGE subsets. \mathbf{A}_{iii} , Pie chart indicating the percentages of cells recovered across the interneuron subtypes from cortex and the hippocampus. **B**, Violin plot showing the distribution of expression levels of well-known representative cell-type-enriched marker genes across the 11 MGE subtypes.

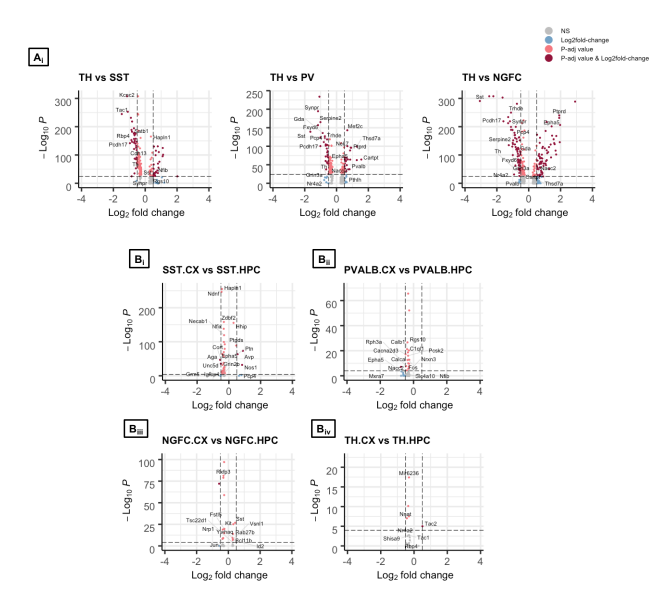


Figure 11: (Figure1, Supp.4) | MGE subtype differences between cortex and hippocampus Volcano plot representing the $-\log 10$ False Discovery Rate (FDR) versus $\log 2$ fold change (FC) between A, TH-expressing MGE subsets and the remaining MGE subset SST, PVALB and NGFC; B, Differential expression of the cardinal MGE classes between cortex and hippocampus, at a fold change ≥ 0.5 and FDR <10e-5.

- 799 URL https://dx.doi.org/10.1038/nrn3969
- 800 [7] T. Klausberger, P. Somogyi (2008).
- [8] G. Akgül, C. J. McBain, Diverse roles for ionotropic glutamate receptors on inhibitory interneurons in developing and adult brain, The Journal of Physiology 594 (19) (2016) 5471–5490.
 doi:10.1113/jp271764.
- 804 URL https://dx.doi.org/10.1113/jp271764
- R. Chittajallu, J. C. Wester, M. T. Craig, E. Barksdale, X. Q. Yuan, G. Akgül, C. Fang, D. Collins,
 S. Hunt, K. A. Pelkey, C. J. McBain, Afferent specific role of NMDA receptors for the circuit integration

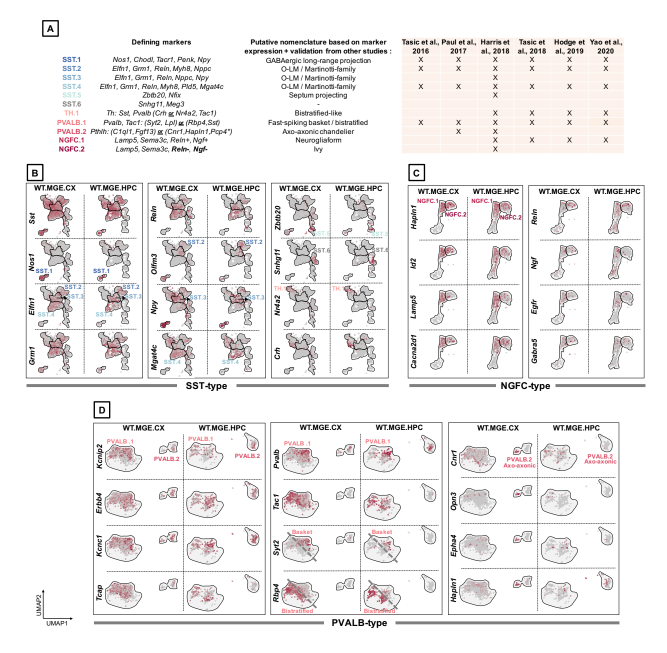


Figure 12: (Figure 1, Supp.5) | MGE subtype annotation based on marker expression A, Table indicating the subype-defining marker genes observed in the present study and their descriptions in the previous scRNAseq datasets (*indicates the genes expressed in the cortex-exclusive PVALB.2 subcluster). Representative UMAP plots of MGE subtype-enriched genes in B, SST subclusters, C, NGFC subclusters, and D, PVALB subclusters.

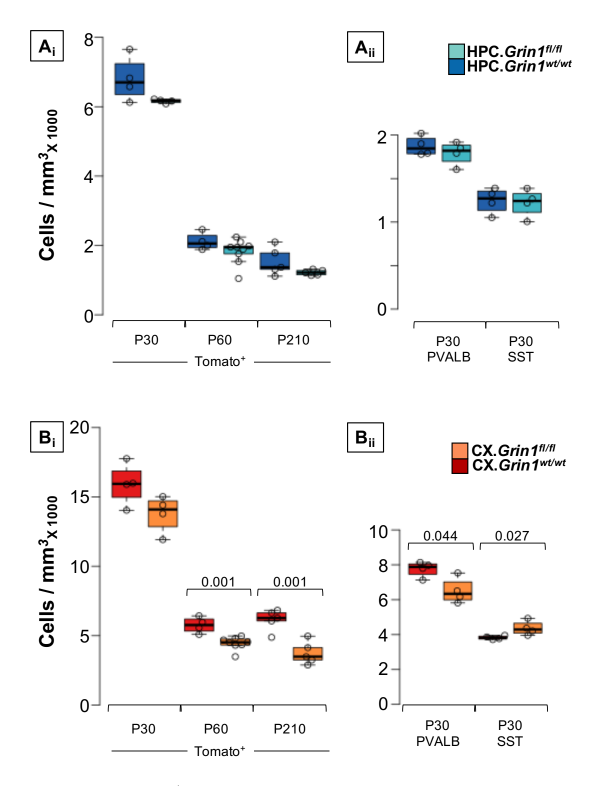
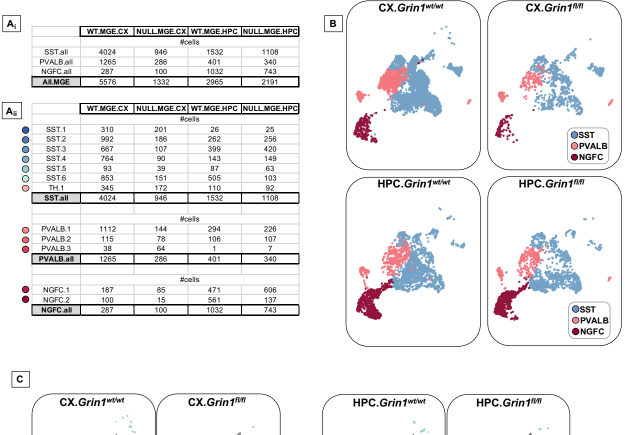


Figure 13: Figure2, Supp.1 | MGE subtype abundances upon *Grin1*-ablation A, Boxplots indicating the cell counts of hippocampal MGE-derived interneurons expressing (A_i) Ai14/tdTomato, (A_{ii}) PV, SST immunostaining from *Grin1*^{wt/wt} and *Grin1*^{fl/fl} B, Boxplots indicating the cell counts of cortical MGE-derived interneurons expressing (B_i) Ai14/tdTomato, and (B_{ii}) PV, SST immunostaining from *Grin1*^{wt/wt} and *Grin1*^{fl/fl}. t-test for statistical analysis.



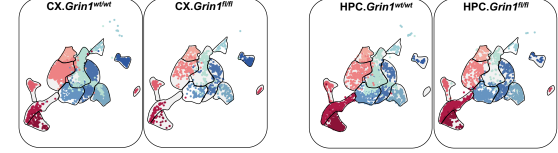


Figure 14: Figure 2, Supp.2 | scRNAseq differential recoveries of MGEsubtypes

 \mathbf{A}_{i} , Number of cells recovered across cardinal subtypes SST, PV and NGFC. \mathbf{A}_{ii} , Number of cells recovered within the subtypes of PV / SST/ NGFC. **B**, UMAP representation colored by cardinal MGE-derived interneuron subtypes SST, PVALB and NGFC, highlighting the differential enrichments of cells **C**, Representative UMAP plots indicating the granularity among PV/SST/NGFC subtypes between both brain regions and both genotypes.

- 807
 of hippocampal neurogliaform cells, Nature Communications 8 (1) (2017) 152–152. doi:10.1038/s41467

 808
 017-00218-y.
- 809 URL https://dx.doi.org/10.1038/s41467-017-00218-y
- [10] J. H. Cornford, M. S. Mercier, M. Leite, V. Magloire, M. Häusser, D. M. Kullmann, Dendritic
 NMDA receptors in parvalbumin neurons enable strong and stable neuronal assemblies (2019).
 doi:10.7554/elife.49872.
- 813 URL https://dx.doi.org/10.7554/elife.49872
- 814 [11] N. V. D. M. García, R. Priya, S. N. Tuncdemir, G. Fishell, T. Karayannis, Sensory inputs control

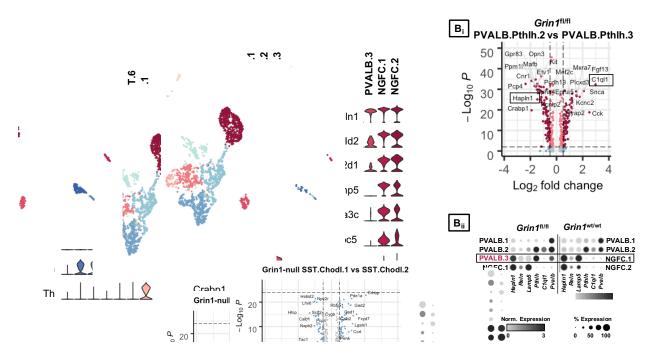
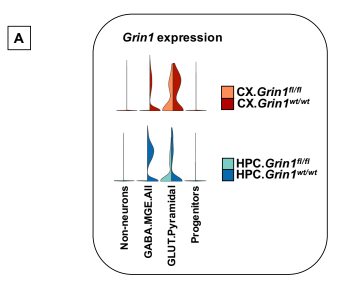


Figure 15: Figure2, Supp.3 | Select marker gene expression across the subtypes of merged MGE-derived interneurons from $Grin1^{wt/wt}$ and $Grin1^{fl/fl}$ A, Violin plot showing the distribution of expression levels of well-known representative cell-type-enriched marker genes across the MGE subtypes. B_i, -log10 False Discovery Rate (FDR) versus log2 fold change (FC) between *Pthlh*-PVALB.2 and *Pthlh*-PVALB.3 at a fold change ≥ 0.5 and FDR <10e-3. B_{ii}, Dot plots representing the normalized expressions of NGFC marker genes mis expressed in *Pthlh*-PVALB.3 upon *Grin1*-ablation.

- the integration of neurogliaform interneurons into cortical circuits, Nature Neuroscience 18 (3) (2015)
- 816 393–401. doi:10.1038/nn.3946.
- 817 URL https://dx.doi.org/10.1038/nn.3946
- [12] J. A. Matta, K. A. Pelkey, M. T. Craig, R. Chittajallu, B. W. Jeffries, C. J. McBain, Developmental
 origin dictates interneuron AMPA and NMDA receptor subunit composition and plasticity, Nature
 Neuroscience 16 (8) (2013) 1032–1041. doi:10.1038/nn.3459.
- 821 URL https://dx.doi.org/10.1038/nn.3459
- [13] R. Luján, R. Shigemoto, G. López-Bendito, Glutamate and GABA receptor signalling in the developing brain, Neuroscience 130 (3) (2005) 567–580. doi:10.1016/j.neuroscience.2004.09.042.
- ⁸²⁴ URL https://dx.doi.org/10.1016/j.neuroscience.2004.09.042
- [14] J. B. Manent, Glutamate Acting on AMPA But Not NMDA Receptors Modulates the Migration of Hip pocampal Interneurons, Journal of Neuroscience 26 (22) (2006) 5901–5909. doi:10.1523/jneurosci.1033 06.2006.
- URL https://dx.doi.org/10.1523/jneurosci.1033-06.2006
- [15] J. M. Soria, Receptor-activated Calcium Signals in Tangentially Migrating Cortical Cells, Cerebral
 Cortex 12 (8) (2002) 831–839. doi:10.1093/cercor/12.8.831.
- uRL https://dx.doi.org/10.1093/cercor/12.8.831
- [16] E. Hanson, M. Armbruster, L. A. Lau, M. E. Sommer, Z.-J. Klaft, S. A. Swanger, S. F. Traynelis, S. J.
 Moss, F. Noubary, J. Chadchankar, C. G. Dulla, Tonic Activation of GluN2C/GluN2D-Containing
 NMDA Receptors by Ambient Glutamate Facilitates Cortical Interneuron Maturation, The Journal
- of Neuroscience 39 (19) (2019) 3611–3626. doi:10.1523/jneurosci.1392-18.2019.
- 836 URL https://dx.doi.org/10.1523/jneurosci.1392-18.2019





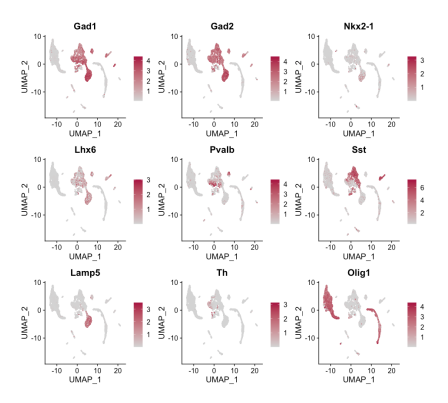


Figure 16: Figure 2, Supp.4 | Expression of control genes in the MGE subtypes subsequent to Grin1-ablation

 \mathbf{A}_i , Split-violin plot from both genotypes indicating the expression of *Grin1* in the MGE-derived interneurons, pyramidal neurons and non-neurons. **B**, Representative UMAP plots of cardinal MGE markers genes.

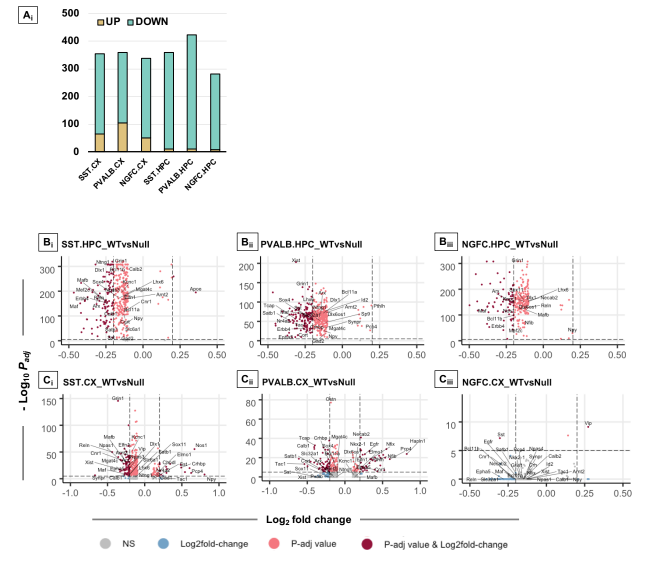


Figure 17: (Figure 3, Supp.1) | Differential gene expression in the MGE subtypes subsequent to Grin1-ablation A_i , Split-violin plot from both genotypes indicating the expression of Grin1 in the MGE-derived interneurons, pyramidal neurons and non-neurons. A_{ii} , Bar plot denoting the number of genes up/downregulated in the cortical and hippocampal MGE clusters. Volcano plot representing the $-\log 10$ False Discovery Rate (FDR) versus $\log 2$ fold change (FC) between **B**, hippocampal and **C**, cortical MGE cardinal clusters upon Grin1-loss, at a fold change ≥ 0.2 and FDR <10e-6.

- [17] W. Kelsch, Z. Li, S. Wieland, O. Senkov, A. Herb, C. Gongrich, H. Monyer, GluN2B-Containing
 NMDA Receptors Promote Glutamate Synapse Development in Hippocampal Interneurons, Journal
 of Neuroscience 34 (48) (2014) 16022–16030. doi:10.1523/jneurosci.1210-14.2014.
- 840 URL https://dx.doi.org/10.1523/jneurosci.1210-14.2014
- [18] G. Akgül, C. J. McBain, AMPA receptor deletion in developing MGE-derived hippocampal interneu rons causes a redistribution of excitatory synapses and attenuates postnatal network oscillatory ac tivity, Scientific Reports 10 (1) (2020) 1333–1333. doi:10.1038/s41598-020-58068-6.
- URL https://dx.doi.org/10.1038/s41598-020-58068-6

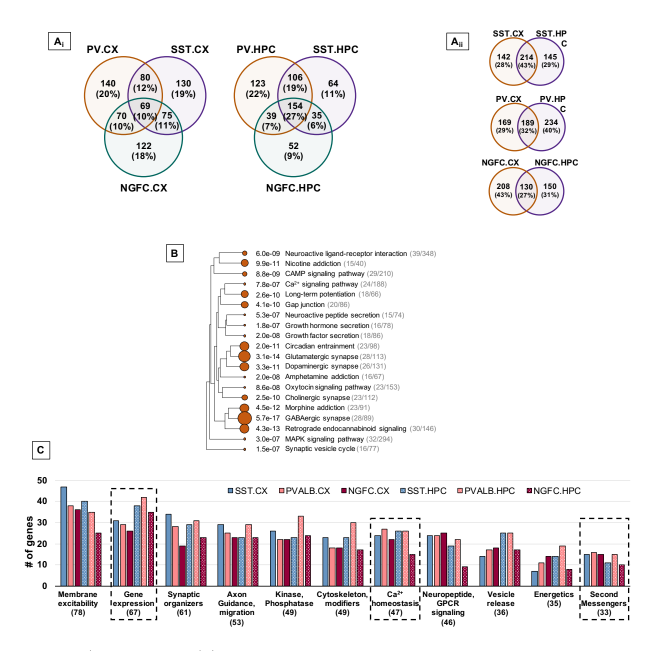
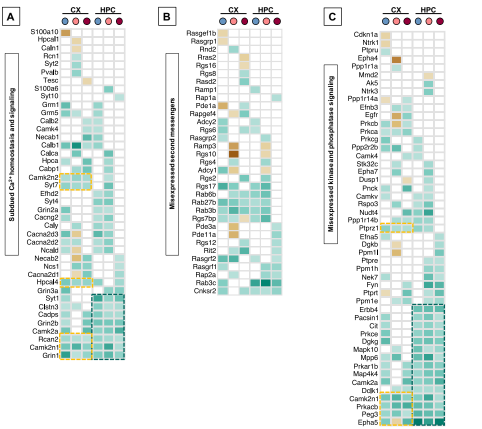


Figure 18: (Figure3, Supp.2) | Molecular pathways differentially expressed in MGE subtypes A_i , Venn-diagrams indicating the percentages of DEGs common within MGE subtypes from cortex or hippocampus. A_i , Venn-diagrams indicating the percentages of DEGs common within MGE subtypes from cortex and hippocampus. B, Hierarchical clustering tree summarizing the correlation among significant pathways enriched among the DEGs. Pathways with many shared genes are clustered together. Bigger dots indicate more significant P-values. C, Bar plot showing the classification of molecular functions of the DEGs, across the MGE subtypes. Total number of DEGs in the particular molecular class indicated in parentheses.



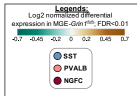


Figure 19: (Figure 4, Supp.1) | Differential expression of intracellular signaling cascades across subtypes upon *Grin1* -ablation Heatmap of log2 FC of significant DEGs in cortical and hippocampal MGE cardinal subtypes, showing a subset of **A**, genes regulating intracellular Ca^{2+} homeostasis and Ca^{2+} binding proteins; **B**, notable second messengers; and, **C**, Ca^{2+} dependent / activated kinases and phosphatases.

- [19] D. Bortone, F. Polleux, KCC2 Expression Promotes the Termination of Cortical Interneuron Migration in a Voltage-Sensitive Calcium-Dependent Manner, Neuron 62 (1) (2009) 53-71.
 doi:10.1016/j.neuron.2009.01.034.
- 848 URL https://dx.doi.org/10.1016/j.neuron.2009.01.034
- [20] C. L. Magueresse, H. Monyer, GABAergic Interneurons Shape the Functional Maturation of the Cortex, Neuron 77 (3) (2013) 388–405. doi:10.1016/j.neuron.2013.01.011.
 - URL https://dx.doi.org/10.1016/j.neuron.2013.01.011

851

- [21] N. V. D. M. García, T. Karayannis, G. Fishell, Neuronal activity is required for the development of specific cortical interneuron subtypes, Nature 472 (7343) (2011) 351–355. doi:10.1038/nature09865.
 URL https://dx.doi.org/10.1038/nature09865
- [22] M. Yozu, H. Tabata, N. König, K. Nakajima, Migratory Behavior of Presumptive Interneurons Is
 Affected by AMPA Receptor Activation in Slice Cultures of Embryonic Mouse Neocortex, Developmental Neuroscience 30 (1-3) (2008) 105–116. doi:10.1159/000109856.
- 858 URL https://dx.doi.org/10.1159/000109856
- [23] O. Marín, Interneuron dysfunction in psychiatric disorders, Nat. Rev. Neurosci 13 (2012) 107–120.
- [24] K. Nakazawa, K. Sapkota, The origin of NMDA receptor hypofunction in schizophrenia, Pharmacology
 & Therapeutics 205 (2020) 107426–107426. doi:10.1016/j.pharmthera.2019.107426.

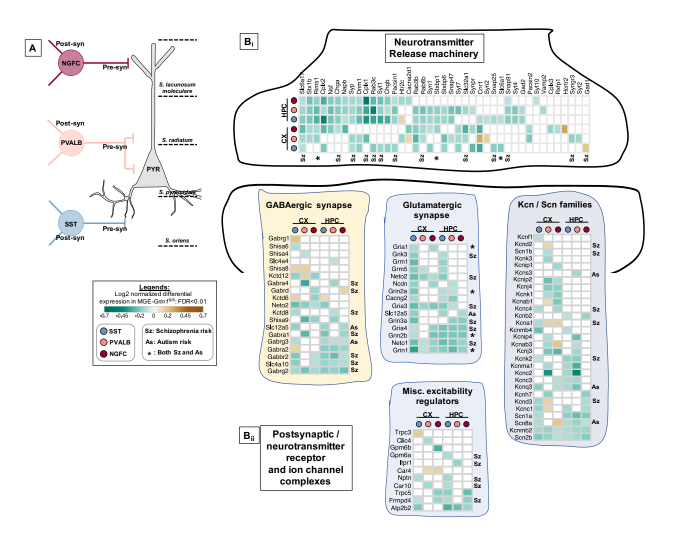


Figure 20: (Figure 6, Supp.1) | Aberrant NMDAR-signaling result in misexpression of regulators of membrane excitability that are high-risk Sz genes A, Schema representing the field of hippocampal pyramidal cell innervated by the interneurons. Heatmap of log2 FC of significant DEGs in cortical and hippocampal MGE cardinal subtypes, showing a subset of \mathbf{B}_{i} , Neurotransmitter release machinery. \mathbf{B}_{ii} , Postsynaptic GABA / Glutamate receptor complexes, *Kcn-, Scn-* ion channel complexes and other miscellaneous regulators of excitability.

- ⁸⁶² URL https://dx.doi.org/10.1016/j.pharmthera.2019.107426
- [25] B. Rico, O. Marín, Neuregulin signaling, cortical circuitry development and schizophrenia, Current
 Opinion in Genetics & Development 21 (3) (2011) 262–270. doi:10.1016/j.gde.2010.12.010.
- 865 URL https://dx.doi.org/10.1016/j.gde.2010.12.010
- [26] A. K. Malhotra, D. A. Pinals, C. M. Adler, I. Elman, A. Clifton, D. Pickar, A. Breier, Ketamineinduced exacerbation of psychotic symptoms and cognitive impairment in neuroleptic-free schizophrenics, Neuropsychopharmacology 17 (1997) 141–150.
- [27] H. Homayoun, B. Moghaddam, NMDA Receptor Hypofunction Produces Opposite Effects on Pre frontal Cortex Interneurons and Pyramidal Neurons, Journal of Neuroscience 27 (43) (2007) 11496–
 11500. doi:10.1523/jneurosci.2213-07.2007.
- URL https://dx.doi.org/10.1523/jneurosci.2213-07.2007
- [28] R. Nguyen, M. D. Morrissey, V. Mahadevan, J. D. Cajanding, M. A. Woodin, J. S. Yeomans,

- K. Takehara-Nishiuchi, J. C. Kim, Parvalbumin and GAD65 Interneuron Inhibition in the Ventral
 Hippocampus Induces Distinct Behavioral Deficits Relevant to Schizophrenia, Journal of Neuroscience
- 876 34 (45) (2014) 14948–14960. doi:10.1523/jneurosci.2204-14.2014.
- 877 URL https://dx.doi.org/10.1523/jneurosci.2204-14.2014
- [29] J. E. Belforte, V. Zsiros, E. R. Sklar, Z. Jiang, G. Yu, Y. Li, E. M. Quinlan, K. Nakazawa, Postnatal
 NMDA receptor ablation in corticolimbic interneurons confers schizophrenia-like phenotypes, Nature
 Neuroscience 13 (1) (2010) 76–83. doi:10.1038/nn.2447.
- 881 URL https://dx.doi.org/10.1038/nn.2447
- [30] T. Korotkova, E. C. Fuchs, A. Ponomarenko, J. von Engelhardt, H. Monyer, NMDA Receptor Ablation
 on Parvalbumin-Positive Interneurons Impairs Hippocampal Synchrony, Spatial Representations, and
 Working Memory, Neuron 68 (3) (2010) 557–569. doi:10.1016/j.neuron.2010.09.017.
- 885 URL https://dx.doi.org/10.1016/j.neuron.2010.09.017
- [31] V. M. Tatard-Leitman, C. R. Jutzeler, J. Suh, J. A. Saunders, E. N. Billingslea, S. Morita, R. White,
 R. E. Featherstone, R. Ray, P. I. Ortinski, A. Banerjee, M. J. Gandal, R. Lin, A. Alexandrescu,
 Y. Liang, R. E. Gur, K. E. Borgmann-Winter, G. C. Carlson, C.-G. Hahn, S. J. Siegel, Pyramidal
 Cell Selective Ablation of N-Methyl-D-Aspartate Receptor 1 Causes Increase in Cellular and Network
 Excitability, Biological Psychiatry 77 (6) (2015) 556–568. doi:10.1016/j.biopsych.2014.06.026.
 URL https://dx.doi.org/10.1016/j.biopsych.2014.06.026
- [32] A. R. Mohn, R. R. Gainetdinov, M. G. Caron, B. H. Koller, Mice with Reduced NMDA Receptor
 Expression Display Behaviors Related to Schizophrenia, Cell 98 (4) (1999) 427–436. doi:10.1016/s00928674(00)81972-8.
- URL https://dx.doi.org/10.1016/s0092-8674(00)81972-8
- [33] A. M. Bygrave, K. Kilonzo, D. M. Kullmann, D. M. Bannerman, D. Kätzel, Can N-Methyl-D-Aspartate Receptor Hypofunction in Schizophrenia Be Localized to an Individual Cell Type?, Frontiers in Psychiatry 10 (2019) 835–835. doi:10.3389/fpsyt.2019.00835.
- 899 URL https://dx.doi.org/10.3389/fpsyt.2019.00835
- [34] C. Mayer, C. Hafemeister, R. C. Bandler, R. Machold, R. B. Brito, X. Jaglin, K. Allaway, A. Butler, G. Fishell, R. Satija, Developmental diversification of cortical inhibitory interneurons, Nature 555 (7697) (2018) 457-462. doi:10.1038/nature25999.
 UBL https://dx.doi.org/10.1038/nature25909.
- 903 URL https://dx.doi.org/10.1038/nature25999
- [35] R. Priya, M. F. Paredes, T. Karayannis, N. Yusuf, X. Liu, X. Jaglin, I. Graef, A. Alvarez-Buylla,
 G. Fishell, Activity Regulates Cell Death within Cortical Interneurons through a CalcineurinDependent Mechanism, Cell Reports 22 (7) (2018) 1695–1709. doi:10.1016/j.celrep.2018.01.007.
 URL https://dx.doi.org/10.1016/j.celrep.2018.01.007
- [36] F. K. Wong, K. Bercsenyi, V. Sreenivasan, A. Portalés, M. Fernández-Otero, O. Marín, Pyramidal
 cell regulation of interneuron survival sculpts cortical networks, Nature 557 (7707) (2018) 668–673.
 doi:10.1038/s41586-018-0139-6.
- [37] H. Bading, D. Ginty, M. Greenberg, Regulation of gene expression in hippocampal neurons by distinct calcium signaling pathways, Science 260 (5105) (1993) 181–186. doi:10.1126/science.8097060.
 URL https://dx.doi.org/10.1126/science.8097060
- [38] H. Bading, M. M. Segal, N. J. Sucher, H. Dudek, ~. Lipton, S. A. Greenberg, M. E (1995).
- [39] E. L. Yap, M. E. Greenberg, Activity-Regulated Transcription: Bridging the Gap between Neural
 Activity and Behavior (2018).
- 917 URL https://doi.org/10.1016/j.neuron.2018.10.013
- [40] S. Cull-Candy, S. Brickley, M. Farrant, NMDA receptor subunits: diversity, development and disease,
 Current Opinion in Neurobiology 11 (3) (2001) 327–335. doi:10.1016/s0959-4388(00)00215-4.
- 920 URL https://dx.doi.org/10.1016/s0959-4388(00)00215-4
- 921 [41] H. Komuro, P. Rakic (1993).
- [42] Q. Xu, M. Tam, S. A. Anderson, Fate mapping Nkx2.1-lineage cells in the mouse telencephalon, The
 Journal of Comparative Neurology 506 (1) (2008) 16–29. doi:10.1002/cne.21529.
- 924 URL https://dx.doi.org/10.1002/cne.21529

- [43] K. D. Harris, H. Hochgerner, N. G. Skene, L. Magno, L. Katona, C. B. Gonzales, P. Somogyi,
 N. Kessaris, S. Linnarsson, J. Hjerling-Leffler, Classes and continua of hippocampal CA1 inhibitory
 neurons revealed by single-cell transcriptomics, PLOS Biology 16 (6) (2018) e2006387–e2006387.
 doi:10.1371/journal.pbio.2006387.
- 929 URL https://dx.doi.org/10.1371/journal.pbio.2006387
- [44] A. B. Muñoz-Manchado, C. B. Gonzales, A. Zeisel, H. Munguba, B. Bekkouche, N. G. Skene, P. Lönnerberg, J. Ryge, K. D. Harris, S. Linnarsson, J. Hjerling-Leffler, Diversity of Interneurons in the Dorsal Striatum Revealed by Single-Cell RNA Sequencing and PatchSeq, Cell Reports 24 (8) (2018)
 2179–2190.e7. doi:10.1016/j.celrep.2018.07.053.
- 934 URL https://dx.doi.org/10.1016/j.celrep.2018.07.053
- [45] Y. Tanaka, Y. Tanaka, T. Furuta, Y. Yanagawa, T. Kaneko, The effects of cutting solutions on the viability of GABAergic interneurons in cerebral cortical slices of adult mice, Journal of Neuroscience Methods 171 (1) (2008) 118–125. doi:10.1016/j.jneumeth.2008.02.021.
- 938 URL https://dx.doi.org/10.1016/j.jneumeth.2008.02.021
- [46] Y. E. Wu, L. Pan, Y. Zuo, X. Li, W. Hong, Detecting Activated Cell Populations Using Single-Cell
 RNA-Seq, Neuron 96 (2017) 1–27.
- [47] A. Butler, P. Hoffman, P. Smibert, E. Papalexi, R. Satija, Integrating single-cell transcriptomic data across different conditions, technologies, and species, Nature Biotechnology 36 (5) (2018) 411-420. doi:10.1038/nbt.4096.
- 944 URL https://dx.doi.org/10.1038/nbt.4096
- [48] T. Stuart, A. Butler, P. Hoffman, C. Hafemeister, E. Papalexi, W. M. Mauck, Y. Hao, M. Stoeckius,
 P. Smibert, R. Satija, Comprehensive Integration of Single-Cell Data. Cell (2019).
- [49] A. Paul, M. Crow, R. Raudales, M. He, J. Gillis, Z. J. Huang, Transcriptional Architecture of Synaptic Communication Delineates GABAergic Neuron Identity, Cell 171 (3) (2017) 522–539.e20. doi:10.1016/j.cell.2017.08.032.
- 950 URL https://dx.doi.org/10.1016/j.cell.2017.08.032
- [50] B. Tasic, V. Menon, T. N. T. Nguyen, T. T. K. Kim, T. Jarsky, Z. Yao, B. B. Levi, L. T. Gray, S. A.
 Sorensen, T. Dolbeare, Adult mouse cortical cell taxonomy revealed by single cell transcriptomics, Nat. Neurosci. advance on (2016) 1–37.
- [51] B. Tasic, V. Menon, T. N. T. Nguyen, T. T. K. Kim, T. Jarsky, Z. Yao, B. B. Levi, L. T. Gray,
 S. A. Sorensen, T. Dolbeare, D. Bertagnolli, J. Goldy, N. Shapovalova, S. Parry, C. C. Lee, K. Smith,
 A. Bernard, L. Madisen, S. M. Sunkin, M. Hawrylycz, C. Koch, H. Zeng, Z. Yao, C. C. Lee, N. Shapovalova, S. Parry, L. Madisen, S. M. Sunkin, M. Hawrylycz, C. Koch, H. Zeng, Adult mouse cortical cell
 taxonomy revealed by single cell transcriptomics, Nature Neuroscience advance on (January) (2016)
 1–37. doi:10.1038/nn.4216.
- [52] J. Winterer, D. Lukacsovich, L. Que, A. M. Sartori, W. Luo, C. Földy, Single-cell RNA-Seq characterization of anatomically identified OLM interneurons in different transgenic mouse lines, European Journal of Neuroscience 50 (11) (2019) 3750–3771. doi:10.1111/ejn.14549.
- 963 URL https://dx.doi.org/10.1111/ejn.14549
- [53] B. Tasic, Z. Yao, L. T. Graybuck, K. A. Smith, T. N. Nguyen, D. Bertagnolli, J. Goldy, E. Garren,
 M. N. Economo, S. Viswanathan, Shared and distinct transcriptomic cell types across neocortical
 areas, Nature 563 (2018) 72–78.
- [54] E. Favuzzi, R. Deogracias, A. Marques-Smith, P. Maeso, J. Jezequel, D. Exposito-Alonso, M. Balia,
 T. Kroon, A. J. Hinojosa, E. F. Maraver, B. Rico, Distinct molecular programs regulate synapse specificity in cortical inhibitory circuits, Science 363 (6425) (2019) 413–417. doi:10.1126/science.aau8977.
- [55] R. Priya, M. F. Paredes, T. Karayannis, N. Yusuf, X. Liu, X. Jaglin, I. Graef, A. Alvarez-Buylla,
 G. Fishell, Activity Regulates Cell Death within Cortical Interneurons through a CalcineurinDependent Mechanism., Cell reports 22 (7) (2018) 1695–1709. doi:10.1016/j.celrep.2018.01.007.
- [56] S. J. Butt, V. H. Sousa, M. V. Fuccillo, J. Hjerling-Leffler, G. Miyoshi, S. Kimura, G. Fishell, The Requirement of Nkx2-1 in the Temporal Specification of Cortical Interneuron Subtypes, Neuron 59 (5)
 (2002) 720, 720, 121 (10, 1016)

- URL https://dx.doi.org/10.1016/j.neuron.2008.07.031 976
- [57] D. J. Laurie, P. H. Seeburg, Regional and developmental heterogeneity in splicing of the rat brain 977 NMDAR1 mRNA, The Journal of Neuroscience 14 (5) (1994) 3180–3194. doi:10.1523/jneurosci.14-978 979 05-03180.1994.
- URL https://dx.doi.org/10.1523/jneurosci.14-05-03180.1994 980
- [58] H. Monyer, N. Burnashev, D. J. Laurie, B. Sakmann, P. H. Seeburg, Developmental and regional 981 expression in the rat brain and functional properties of four NMDA receptors, Neuron 12 (3) (1994) 982 529-540. doi:10.1016/0896-6273(94)90210-0. 983
- URL https://dx.doi.org/10.1016/0896-6273(94)90210-0 984
- [59] Z. J. Huang, A. Paul, The diversity of GABAergic neurons and neural communication elements, Nature 985 Reviews Neuroscience 20 (9) (2019) 563-572. doi:10.1038/s41583-019-0195-4. 986 987
 - URL https://dx.doi.org/10.1038/s41583-019-0195-4
- [60] O. Hobert, I. Carrera, N. Stefanakis, The molecular and gene regulatory signature of a neuron, Trends 988 in Neurosciences 33 (10) (2010) 435–445. doi:10.1016/j.tins.2010.05.006. 980
- URL https://dx.doi.org/10.1016/j.tins.2010.05.006 990
- [61] K. Y. Kwan, N. Sestan, E. S. Anton, Transcriptional co-regulation of neuronal migration and laminar 991 identity in the neocortex, Development 139 (9) (2012) 1535–1546. doi:10.1242/dev.069963. 992 URL https://dx.doi.org/10.1242/dev.069963 993
- [62] R. Janky, A. Verfaillie, H. Imrichová, B. V. de Sande, L. Standaert, V. Christiaens, G. Hulselmans, 994 K. Herten, M. N. Sanchez, D. Potier, D. Svetlichnyv, Z. K. Atak, M. Fiers, J.-C. Marine, S. Aerts, 995 iRegulon: From a Gene List to a Gene Regulatory Network Using Large Motif and Track Collections, 996 PLoS Computational Biology 10 (7) (2014) e1003731-e1003731. doi:10.1371/journal.pcbi.1003731. 997 URL https://dx.doi.org/10.1371/journal.pcbi.1003731 998
- [63] S. J. Otto, S. R. McCorkle, J. Hover, C. Conaco, J. J. Han, S. Impey, G. S. Yochum, J. J. Dunn, R. H. 999 Goodman, G. Mandel, A New Binding Motif for the Transcriptional Repressor REST Uncovers Large 1000 Gene Networks Devoted to Neuronal Functions, Journal of Neuroscience 27 (25) (2007) 6729–6739. 1001 doi:10.1523/jneurosci.0091-07.2007. 1002
- URL https://dx.doi.org/10.1523/jneurosci.0091-07.2007 1003
- [64] A. Rodenas-Ruano, A. E. Chávez, M. J. Cossio, P. E. Castillo, R. S. Zukin, REST-dependent epigenetic 1004 remodeling promotes the developmental switch in synaptic NMDA receptors, Nature Neuroscience 1005 15 (10) (2012) 1382–1390. doi:10.1038/nn.3214. 1006
- URL https://dx.doi.org/10.1038/nn.3214 1007
- [65] K. Nikouei, A. B. Muñoz-Manchado, J. Hjerling-Leffler, BCL11B/CTIP2 is highly expressed in 1008 GABAergic interneurons of the mouse somatosensory cortex, Journal of Chemical Neuroanatomy 1009 71 (2016) 1-5. doi:10.1016/j.jchemneu.2015.12.004. 1010
- URL https://dx.doi.org/10.1016/j.jchemneu.2015.12.004 1011
- [66] M. Erburu, I. Muñoz-Cobo, T. Diaz-Perdigon, P. Mellini, T. Suzuki, E. Puerta, R. M. 1012 Tordera, SIRT2 inhibition modulate glutamate and serotonin systems in the prefrontal 1013 cortex and induces antidepressant-like action, Neuropharmacology 117 (2017) 195–208. 1014 doi:10.1016/j.neuropharm.2017.01.033. 1015
- URL https://dx.doi.org/10.1016/j.neuropharm.2017.01.033 1016
- [67] M. Watroba, D. Szukiewicz, The role of sirtuins in aging and age-related diseases, Advances in Medical 1017 Sciences 61 (1) (2016) 52–62. doi:10.1016/j.advms.2015.09.003. 1018
- URL https://dx.doi.org/10.1016/j.advms.2015.09.003 1019
- [68] N. Sharma, E. A. Pollina, M. A. Nagy, E.-L. Yap, F. A. DiBiase, S. Hrvatin, L. Hu, C. Lin, M. E. 1020 Greenberg, ARNT2 Tunes Activity-Dependent Gene Expression through NCoR2-Mediated Repression 1021 and NPAS4-Mediated Activation, Neuron 102 (2) (2019) 390-406.e9. doi:10.1016/j.neuron.2019.02.007. 1022 URL https://dx.doi.org/10.1016/j.neuron.2019.02.007 1023
- [69] Y. Wu, Y. G. Yao, X. J. Luo, SZDB: A Database for Schizophrenia Genetic Research, Schizophr. Bull 1024 43 (2017) 459-471. 1025
- [70] S. Foundation (2018). 1026

- [71] A. S. Cristino, S. M. Williams, Z. Hawi, J. Y. An, M. A. Bellgrove, C. E. Schwartz, L. da F Costa, 1027 C. Claudianos, Neurodevelopmental and neuropsychiatric disorders represent an interconnected molec-1028 ular system, Molecular Psychiatry 19 (3) (2014) 294–301. doi:10.1038/mp.2013.16. 1029 1030
 - URL https://dx.doi.org/10.1038/mp.2013.16
- [72]J. Liu, M. Li, X. J. Luo, B. Su, Systems-level analysis of risk genes reveals the modular nature of 1031 schizophrenia, Schizophr. Res 201 (2018) 261-269. 1032
- [73] K. J. Mitchell, The genetics of neurodevelopmental disease, Curr. Opin. Neurobiol 21 (2011) 197–203. 1033
- [74] Y. Sakai, C. A. Shaw, B. C. Dawson, D. V. Dugas, Z. Al-Mohtaseb, D. E. Hill, H. Y. Zoghbi, Protein 1034 Interactome Reveals Converging Molecular Pathways Among Autism Disorders, Science Translational 1035 Medicine 3 (86) (2011) 86ra49-86ra49. doi:10.1126/scitranslmed.3002166. 1036 URL https://dx.doi.org/10.1126/scitranslmed.3002166 1037
- [75] E. Favuzzi, R. Deogracias, A. Marques-Smith, P. Maeso, J. Jezequel, D. Exposito-Alonso, M. Balia, 1038 T. Kroon, A. J. Hinojosa, E. F. Maraver, B. Rico, Distinct molecular programs regulate synapse speci-1039 ficity in cortical inhibitory circuits, Science 363 (6425) (2019) 413–417. doi:10.1126/science.aau8977. 1040
- URL https://dx.doi.org/10.1126/science.aau8977 1041
- [76] R. Dolmetsch, Excitation-Transcription Coupling: Signaling by Ion Channels to the Nucleus, Science 1042 Signaling 2003 (166) (2003) pe4-pe4. doi:10.1126/stke.2003.166.pe4. 1043
- URL https://dx.doi.org/10.1126/stke.2003.166.pe4 1044
- [77] S. M. Cohen, H. Ma, K. V. Kuchibhotla, B. O. Watson, G. Buzsáki, R. C. Froemke, R. W. 1045 Tsien, Excitation-Transcription Coupling in Parvalbumin-Positive Interneurons Employs a Novel 1046 CaM Kinase-Dependent Pathway Distinct from Excitatory Neurons, Neuron 90 (2) (2016) 292–307. 1047 doi:10.1016/j.neuron.2016.03.001. 1048
- URL https://dx.doi.org/10.1016/j.neuron.2016.03.001 1049
- [78] N. Flames, R. Pla, D. M. Gelman, J. L. R. Rubenstein, L. Puelles, O. Marin, Delineation of Multiple 1050 Subpallial Progenitor Domains by the Combinatorial Expression of Transcriptional Codes, Journal of 1051 Neuroscience 27 (36) (2007) 9682–9695. doi:10.1523/jneurosci.2750-07.2007. 1052 URL https://dx.doi.org/10.1523/jneurosci.2750-07.2007 1053
- [79] L. Lim, D. Mi, A. Llorca, O. Marín, Development and Functional Diversification of Cortical Interneu-1054 rons, Neuron 100 (2) (2018) 294-313. doi:10.1016/j.neuron.2018.10.009. 1055 1056
 - URL https://dx.doi.org/10.1016/j.neuron.2018.10.009
- [80] N. Dehorter, G. Ciceri, G. Bartolini, L. Lim, I. del Pino, O. Marín, Tuning of fast-spiking interneu-1057 ron properties by an activity-dependent transcriptional switch, Science 349 (6253) (2015) 1216–1220. 1058 doi:10.1126/science.aab3415. 1059
- URL https://dx.doi.org/10.1126/science.aab3415 1060
- [81] P. Flandin, Y. Zhao, D. Vogt, J. Jeong, J. Long, G. Potter, H. Westphal, J. L. R. Rubenstein, Lhx6 and 1061 Lhx8 Coordinately Induce Neuronal Expression of Shh that Controls the Generation of Interneuron 1062 Progenitors, Neuron 70 (5) (2011) 939–950. doi:10.1016/j.neuron.2011.04.020. 1063
- URL https://dx.doi.org/10.1016/j.neuron.2011.04.020 1064
- [82] E. L.-L. Pai, D. Vogt, A. Clemente-Perez, G. L. McKinsey, F. S. Cho, J. S. Hu, M. Wimer, A. Paul, 1065 S. F. Darbandi, R. Pla, T. J. Nowakowski, L. V. Goodrich, J. T. Paz, J. L. Rubenstein, Mafb and 1066 c-Maf Have Prenatal Compensatory and Postnatal Antagonistic Roles in Cortical Interneuron Fate 1067 and Function, Cell Reports 26 (5) (2019) 1157–1173.e5. doi:10.1016/j.celrep.2019.01.031. 1068
- URL https://dx.doi.org/10.1016/j.celrep.2019.01.031 1069
- [83] A. Stanco, R. Pla, D. Vogt, Y. Chen, S. Mandal, J. Walker, R. F. Hunt, S. Lindtner, C. A. 1070 Erdman, A. A. Pieper, S. P. Hamilton, D. Xu, S. C. Baraban, J. L. R. Rubenstein, NPAS1 Re-1071 presses the Generation of Specific Subtypes of Cortical Interneurons, Neuron 84 (5) (2014) 940–953. 1072 doi:10.1016/j.neuron.2014.10.040. 1073
- URL https://dx.doi.org/10.1016/j.neuron.2014.10.040 1074
- [84] Z. Liu, Z. Zhang, S. Lindtner, Z. Li, Z. Xu, S. Wei, Q. Liang, Y. Wen, G. Tao, Y. You, Sp9 Regulates 1075 Medial Ganglionic Eminence-Derived Cortical Interneuron Development, Cereb. Cortex (2018) 1–15. 1076
- [85] R. D. Hodge, T. E. Bakken, J. A. Miller, K. A. Smith, E. R. Barkan, L. T. Graybuck, J. L. Close, 1077

- B. Long, N. Johansen, O. Penn, Conserved cell types with divergent features in human versus mouse cortex, Nature (2019).
- [86] Z. Yao, T. N. Nguyen, C. T. J. Velthoven, Van, J. Goldy, A. E. Sedeno-Cortes, F. Baftizadeh,
 D. Bertagnolli, T. Casper, K. Crichton, S. L. Ding (2020).
- [87] G. Fishell, A. Kepecs, Interneuron Types as Attractors and Controllers (2020). doi:10.1146/annurevneuro-070918-050421.
- 1084 URL https://dx.doi.org/10.1146/annurev-neuro-070918-050421
- [88] W. Andrews, M. Barber, L. R. Hernadez-Miranda, J. Xian, S. Rakic, V. Sundaresan, T. H. Rabbitts, R. Pannell, P. Rabbitts, H. Thompson, L. Erskine, F. Murakami, J. G. Parnavelas, The role of Slit-Robo signaling in the generation, migration and morphological differentiation of cortical interneurons, Developmental Biology 313 (2) (2008) 648–658. doi:10.1016/j.ydbio.2007.10.052.
 URL https://dx.doi.org/10.1016/j.ydbio.2007.10.052
- [89] F. Polleux, K. L. Whitford, P. A. Dijkhuizen, T. Vitalis, A. Ghosh, Control of cortical interneuron migration by neurotrophins and PI3-kinase signaling, Development 129 (2002) 3147–3160.
- [90] O. Marín, A. Yaron, A. Bagri, M. Tessier-Lavigne, J. L. R. Rubenstein, Sorting of Striatal and Cortical
 Interneurons Regulated by Semaphorin-Neuropilin Interactions, Science 293 (2001) 872–875.
- [91] S. Nobrega-Pereira, O. Marin, Transcriptional Control of Neuronal Migration in the Developing Mouse
 Brain, Cerebral Cortex 19 (suppl 1) (2009) i107–i113. doi:10.1093/cercor/bhp044.
 URL https://dx.doi.org/10.1093/cercor/bhp044
- [92] S. Nóbrega-Pereira, N. Kessaris, T. Du, S. Kimura, S. A. Anderson, O. Marín, Postmitotic Nkx2-1
 Controls the Migration of Telencephalic Interneurons by Direct Repression of Guidance Receptors, Neuron 59 (5) (2008) 733-745. doi:10.1016/j.neuron.2008.07.024.
- 1100 URL https://dx.doi.org/10.1016/j.neuron.2008.07.024
- [93] A. Stanco, C. Szekeres, N. Patel, S. Rao, K. Campbell, J. A. Kreidberg, F. Polleux, E. S. Anton, Netrin-1- 3 1 integrin interactions regulate the migration of interneurons through the cortical marginal zone, Proceedings of the National Academy of Sciences 106 (18) (2009) 7595–7600. doi:10.1073/pnas.0811343106.
- 1105 URL https://dx.doi.org/10.1073/pnas.0811343106
- [94] A. Steinecke, C. Gampe, G. Zimmer, J. Rudolph, J. Bolz, EphA/ephrin A reverse signaling promotes the migration of cortical interneurons from the medial ganglionic eminence, Development 141 (2) (2014) 460-471. doi:10.1242/dev.101691.
- 1109 URL https://dx.doi.org/10.1242/dev.101691
- [95] G. Zimmer, P. Garcez, J. Rudolph, R. Niehage, F. Weth, R. Lent, J. Bolz, Ephrin-A5 acts as a repulsive cue for migrating cortical interneurons, European Journal of Neuroscience 28 (1) (2008) 62–73. doi:10.1111/j.1460-9568.2008.06320.x.
- 1113 URL https://dx.doi.org/10.1111/j.1460-9568.2008.06320.x
- [96] Z. Shao, H. Noh, W. B. Kim, P. Ni, C. Nguyen, S. E. Cote, E. Noyes, J. Zhao, T. Parsons, J. M.
 Park, Dysregulated protocadherin-pathway activity as an intrinsic defect in induced pluripotent stem
 cell-derived cortical interneurons from subjects with schizophrenia, Nat. Neurosci 22 (2019) 229–242.
- [97] B. L. Bloodgood, N. Sharma, H. A. Browne, A. Z. Trepman, M. E. Greenberg, The activity-dependent transcription factor NPAS4 regulates domain-specific inhibition, Nature 503 (7474) (2013) 121–125. doi:10.1038/nature12743.
- 1120 URL https://dx.doi.org/10.1038/nature12743
- [98] Y. Lin, B. L. Bloodgood, J. L. Hauser, A. D. Lapan, A. C. Koon, T.-K. Kim, L. S. Hu, A. N. Malik,
 M. E. Greenberg, Activity-dependent regulation of inhibitory synapse development by Npas4, Nature
 455 (7217) (2008) 1198–1204. doi:10.1038/nature07319.
- 1124 URL https://dx.doi.org/10.1038/nature07319
- 1125 [99] X. Sun, Lin, Y. Npas4: Linking Neuronal Activity to Memory, Trends Neurosci 39 (2016) 264–275.
- 1126 [100] I. Spiegel, A. R. Mardinly, H. W. Gabel, J. E. Bazinet, C. H. Couch, C. P. Tzeng, D. A. Harmin, M. E.
- 1127Greenberg, Npas4 Regulates Excitatory-Inhibitory Balance within Neural Circuits through Cell-Type-1128Specific Gene Programs, Cell 157 (5) (2014) 1216–1229. doi:10.1016/j.cell.2014.03.058.

- 1129 URL https://dx.doi.org/10.1016/j.cell.2014.03.058
- [101] B. Ataman, G. L. Boulting, D. A. Harmin, M. G. Yang, M. Baker-Salisbury, E.-L. Yap, A. N. Malik,
 K. Mei, A. A. Rubin, I. Spiegel, E. Durresi, N. Sharma, L. S. Hu, M. Pletikos, E. C. Griffith, J. N.
 Partlow, C. R. Stevens, M. Adli, M. Chahrour, N. Sestan, C. A. Walsh, V. K. Berezovskii, M. S.
 Livingstone, M. E. Greenberg, Evolution of Osteocrin as an activity-regulated factor in the primate
- 1134 brain, Nature 539 (7628) (2016) 242–247. doi:10.1038/nature20111.
- 1135 URL https://dx.doi.org/10.1038/nature20111
- [102] B. Lacar, S. B. Linker, B. N. Jaeger, S. R. Krishnaswami, J. J. Barron, M. J. E. Kelder, S. L. Parylak,
 A. C. M. Paquola, P. Venepally, M. Novotny, Nuclear RNA-seq of single neurons reveals molecular
 signatures of activation, Nat. Commun 7 (2016) 11022–11022.
- [103] C. M. Müller, A. Vlachos, T. Deller, Calcium homeostasis of acutely denervated and lesioned dentate gyrus in organotypic entorhino-hippocampal co-cultures, Cell Calcium 47 (3) (2010) 242–252.
 doi:10.1016/j.ceca.2009.12.006.
- 1142 URL https://dx.doi.org/10.1016/j.ceca.2009.12.006
- [104] W. XiangWei, Y. Jiang, H. Yuan, De novo mutations and rare variants occurring in NMDA receptors, Current Opinion in Physiology 2 (2018) 27–35. doi:10.1016/j.cophys.2017.12.013.
- 1145 URL https://dx.doi.org/10.1016/j.cophys.2017.12.013
- 1146[105] J. C. Masdeu, J. Dalmau, K. F. Berman, NMDA Receptor Internalization by Autoantibodies:1147A Reversible Mechanism Underlying Psychosis?, Trends in Neurosciences 39 (5) (2016) 300–310.1148doi:10.1016/j.tins.2016.02.006.
- 1149 URL https://dx.doi.org/10.1016/j.tins.2016.02.006
- [106] J. Tarabeux, , O. Kebir, J. Gauthier, F. F. Hamdan, L. Xiong, A. Piton, D. Spiegelman, E. Henrion,
 B. Millet, F. Fathalli, R. Joober, J. L. Rapoport, L. E. DeLisi, E. Fombonne, L. Mottron, N. ForgetDubois, M. Boivin, J. L. Michaud, P. Drapeau, R. G. Lafrenière, G. A. Rouleau, M. O. Krebs, Rare
 mutations in N-methyl-D-aspartate glutamate receptors in autism spectrum disorders and schizophrenia, Translational Psychiatry 1 (11) (2011) e55–e55. doi:10.1038/tp.2011.52.
- 1155 URL https://dx.doi.org/10.1038/tp.2011.52
- [107] Y. Yu, Y. Lin, Y. Takasaki, C. Wang, H. Kimura, J. Xing, K. Ishizuka, M. Toyama, I. Kushima,
 D. Mori, Rare loss of function mutations in N-methyl-d-aspartate glutamate receptors and their contributions to schizophrenia susceptibility, Transl. Psychiatry 8 (2018) 12–12.
- [108] W. Chen, C. Shieh, S. A. Swanger, A. Tankovic, M. Au, M. McGuire, M. Tagliati, J. M. Graham,
 S. Madan-Khetarpal, S. F. Traynelis, H. Yuan, T. M. Pierson, GRIN1 mutation associated with
 intellectual disability alters NMDA receptor trafficking and function, Journal of Human Genetics
 62 (6) (2017) 589–597. doi:10.1038/jhg.2017.19.
- 1163 URL https://dx.doi.org/10.1038/jhg.2017.19
- [109] L. L. Gibson, T. A. Pollak, G. Blackman, M. Thornton, N. Moran, A. S. David, The Psychiatric Phenotype of Anti-NMDA Receptor Encephalitis, The Journal of Neuropsychiatry and Clinical Neurosciences 31 (1) (2019) 70–79. doi:10.1176/appi.neuropsych.17120343.
- 1167 URL https://dx.doi.org/10.1176/appi.neuropsych.17120343
- [110] G. L. Carvill, B. M. Regan, S. C. Yendle, B. J. O'Roak, N. Lozovaya, N. Bruneau, N. Burnashev,
 A. Khan, J. Cook, E. Geraghty, L. G. Sadleir, S. J. Turner, M.-H. Tsai, R. Webster, R. Ouvrier, J. A.
 Damiano, S. F. Berkovic, J. Shendure, M. S. Hildebrand, P. Szepetowski, I. E. Scheffer, H. C. Mefford,
 GRIN2A mutations cause epilepsy-aphasia spectrum disorders, Nature Genetics 45 (9) (2013) 1073–
 1076. doi:10.1038/ng.2727.
- 1173 URL https://dx.doi.org/10.1038/ng.2727
- [111] J. R. Lemke, D. Lal, E. M. Reinthaler, I. Steiner, M. Nothnagel, M. Alber, K. Geider, B. Laube,
 M. Schwake, K. Finsterwalder, Mutations in GRIN2A cause idiopathic focal epilepsy with rolandic spikes, Nat. Genet 45 (2013) 1067–1072.
- [112] S. J. Dienel, D. A. Lewis, Alterations in cortical interneurons and cognitive function in schizophrenia, Neurobiology of Disease 131 (2019) 104208–104208. doi:10.1016/j.nbd.2018.06.020.
- 1179 URL https://dx.doi.org/10.1016/j.nbd.2018.06.020

- [113] D. A. Lewis, T. Hashimoto, D. W. Volk, Cortical inhibitory neurons and schizophrenia, Nature Reviews
 Neuroscience 6 (4) (2005) 312–324. doi:10.1038/nrn1648.
- 1182 URL https://dx.doi.org/10.1038/nrn1648
- [114] F. Ali, D. M. Gerhard, K. Sweasy, S. Pothula, C. Pittenger, R. S. Duman, A. C. Kwan, Ketamine disinhibits dendrites and enhances calcium signals in prefrontal dendritic spines, Nature Communications 11 (1) (2020) 72–72. doi:10.1038/s41467-019-13809-8.
- 1186 URL https://dx.doi.org/10.1038/s41467-019-13809-8
- [115] R. Chittajallu, K. Auville, V. Mahadevan, M. Lai, S. Hunt, D. Calvigioni, K. A. Pelkey, K. Zaghloul,
 C. J. Mcbain (2020).
- [116] K. Nakao, V. Jeevakumar, S. Z. Jiang, Y. Fujita, N. B. Diaz, C. A. P. Annan, K. L. E. Jaunarajs, K. Hashimoto, J. E. Belforte, K. Nakazawa (2018).
- 1191 [117] R. C. Team (2013).
- [118] A. Paul, M. Crow, R. Raudales, J. Gillis, Z. J. Huang, Transcriptional Architecture of Synaptic Communication Delineates Cortical GABAergic Neuron Identity, Cell 171 (3) (2017) 522–539.
 doi:10.1016/j.cell.2017.08.032.
- [119] A. Saunders, E. Z. Macosko, A. Wysoker, M. Goldman, F. M. Krienen, H. de Rivera, E. Bien, M. Baum,
 L. Bortolin, S. Wang, A. Goeva, J. Nemesh, N. Kamitaki, S. Brumbaugh, D. Kulp, S. A. McCarroll,
 Molecular Diversity and Specializations among the Cells of the Adult Mouse Brain, Cell 174 (4) (2018)
 1015–1030.e16. doi:10.1016/j.cell.2018.07.028.
- 1199 URL https://dx.doi.org/10.1016/j.cell.2018.07.028
- [120] Z. Yao, T. N. Nguyen, C. T. J. van Velthoven, J. Goldy, A. E. Sedeno-Cortes, F. Baftizadeh, D. Bertagnolli, T. Casper, K. Crichton, S.-L. Ding, O. Fong, E. Garren, A. Glandon, J. Gray, L. T. Graybuck, D. Hirschstein, M. Kroll, K. Lathia, B. Levi, D. McMillen, S. Mok, T. Pham, Q. Ren, C. Rimorin, N. Shapovalova, J. Sulc, S. M. Sunkin, M. Tieu, A. Torkelson, H. Tung, K. Ward, N. Dee, K. A. Smith, B. Tasic, H. Zeng, A taxonomy of transcriptomic cell types across the isocortex and hippocampal formation, bioRxiv (mar 2020). doi:10.1101/2020.03.30.015214.
- [121] G. Finak, A. McDavid, M. Yajima, J. Deng, V. Gersuk, A. K. Shalek, C. K. Slichter, H. W. Miller,
 M. J. McElrath, M. Prlic, P. S. Linsley, R. Gottardo, MAST: a flexible statistical framework for
 assessing transcriptional changes and characterizing heterogeneity in single-cell RNA sequencing data,
 Genome Biology 16 (1) (2015). doi:10.1186/s13059-015-0844-5.
- 1210 URL https://dx.doi.org/10.1186/s13059-015-0844-5
- [122] M. Ximerakis, S. L. Lipnick, B. T. Innes, S. K. Simmons, X. Adiconis, D. Dionne, B. A. Mayweather,
 L. Nguyen, Z. Niziolek, C. Ozek, V. L. Butty, R. Isserlin, S. M. Buchanan, S. S. Levine, A. Regev,
 G. D. Bader, J. Z. Levin, L. L. Rubin, Single-cell transcriptomic profiling of the aging mouse brain,
 Nature Neuroscience 22 (10) (2019) 1696–1708. doi:10.1038/s41593-019-0491-3.
- 1215 URL https://dx.doi.org/10.1038/s41593-019-0491-3
- 1216 [123] B. Kevin (2019). [link].
- 1217 URL Https://Github.Com/Kevinblighe/EnhancedVolcano
- 1218 [124] S. X. Ge, D. Jung, R. Yao, ShinyGO: a graphical gene-set enrichment tool for animals and plants, Bioinformatics 36 (8) (2020) 2628–2629. doi:10.1093/bioinformatics/btz931.
- 1220 URL https://dx.doi.org/10.1093/bioinformatics/btz931
- 1221 [125] K. R. Brown, I. Jurisica, Unequal evolutionary conservation of human protein interactions in interol-1222 ogous networks, Genome Biology 8 (5) (2007) R95–R95. doi:10.1186/gb-2007-8-5-r95.
- 1223 URL https://dx.doi.org/10.1186/gb-2007-8-5-r95
- [126] M. Kotlyar, C. Pastrello, F. Pivetta, A. L. Sardo, C. Cumbaa, H. Li, T. Naranian, Y. Niu, Z. Ding,
 F. Vafaee, F. Broackes-Carter, J. Petschnigg, G. B. Mills, A. Jurisicova, I. Stagljar, R. Maestro,
 I. Jurisica, In silico prediction of physical protein interactions and characterization of interactome
 orphans, Nature Methods 12 (1) (2015) 79–84. doi:10.1038/nmeth.3178.
- 1228 URL https://dx.doi.org/10.1038/nmeth.3178
- 1229 [127] C. Stark, BioGRID: a general repository for interaction datasets, Nucleic Acids Research 34 (90001) 1230 (2006) D535–D539. doi:10.1093/nar/gkj109.

1231 URL https://dx.doi.org/10.1093/nar/gkj109

- [123] V. Mahadevan, C. S. Khademullah, Z. Dargaei, J. Chevrier, P. Uvarov, J. Kwan, R. D. Bagshaw,
 T. Pawson, A. Emili, Y. D. Koninck (2017).
- 1234 [129] M. E. Smoot, K. Ono, J. Ruscheinski, P. L. Wang, T. Ideker, Cytoscape 2.8: new fea-1235 tures for data integration and network visualization, Bioinformatics 27 (3) (2011) 431–432. 1236 doi:10.1093/bioinformatics/btq675.
- 1237 URL https://dx.doi.org/10.1093/bioinformatics/btq675