Title: A single-cell transcriptomic atlas of sensory-dependent gene expression in
 developing mouse visual cortex

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#### 14 Abstract:

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16 Sensory experience drives the refinement and maturation of neural circuits during 17 postnatal brain development through molecular mechanisms that remain to be fully elucidated. One likely mechanism involves the sensory-dependent expression of genes 18 19 that encode direct mediators of circuit remodeling within developing cells. However, while 20 studies in adult systems have begun to uncover crucial roles for sensory-induced genes 21 in modifying circuit connectivity, the gene programs induced by brain cells in response to sensory experience during development remain to be fully characterized. Here, we 22 23 present a single-nucleus RNA-sequencing dataset describing the transcriptional 24 responses of cells in mouse visual cortex to sensory deprivation or sensory stimulation 25 during a developmental window when visual input is necessary for circuit refinement. We 26 sequenced 118,529 individual nuclei across sixteen neuronal and non-neuronal cortical 27 cell types isolated from control, sensory deprived, and sensory stimulated mice, 28 identifying 1,268 unique sensory-induced genes within the developing brain. To 29 demonstrate the utility of this resource, we compared the architecture and ontology of 30 sensory-induced gene programs between cell types, annotated transcriptional induction 31 and repression events based upon RNA velocity, and discovered Neurexin and 32 Neuregulin signaling networks that underlie cell-cell interactions via CellChat. We find that 33 excitatory neurons, especially layer 2/3 pyramidal neurons, are highly sensitive to sensory stimulation, and that the sensory-induced genes in these cells are poised to strengthen 34 synapse-to-nucleus crosstalk by heightening protein serine/threonine kinase activity. 35 36 Altogether, we expect this dataset to significantly broaden our understanding of the 37 molecular mechanisms through which sensory experience shapes neural circuit wiring in 38 the developing brain.

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#### 40 Introduction:

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The precise connectivity of neural circuits in the mammalian brain arises from a convergence of genetic and environmental factors spanning embryonic and postnatal stages of development. Brain circuits are first assembled *in utero* via the formation of an overabundance of nascent synaptic connections between potential neuronal partners, then later remodeled, or *refined*, postnatally through the strengthening of some of these

synapses and the elimination of others<sup>1,2</sup>. The selective retention and maturation of a 47 subset of initially formed synapses equips the brain with an interconnected network of 48 circuits optimized to facilitate neurological function and plasticity across the lifespan. 49 50 Furthermore, impairments in postnatal phases of synaptic remodeling and refinement are 51 increasingly appreciated to contribute to a host of neurodevelopmental conditions such as autism and schizophrenia<sup>3-5</sup>. Thus, elucidating the mechanisms underlying circuit 52 refinement in the early postnatal brain is important from both basic and translational 53 54 perspectives.

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While much emphasis has been placed on defining the intrinsic genetic 56 57 mechanisms that govern embryonic stages of brain development, such as neurogenesis, 58 cell migration, and synapse formation, less is known about how environmental triggers 59 that come online after birth shape the maturation of neural circuits as they emerge. A 60 prime example of environmental cues impacting circuit development can be seen in the role of sensory experience in driving the refinement of neural circuitry within the visual 61 62 system of the mouse, a process that takes place around the third week of life<sup>6</sup>. Specifically, between postnatal days (P)20 and P30, visual experience promotes the 63 structural and functional refinement and maturation of synaptic connections between 64 excitatory thalamocortical neurons within the dorsal lateral geniculate nucleus (dLGN) of 65 the thalamus and their postsynaptic targets in layer 4 of primary visual cortex (V1)<sup>7-10</sup>. 66 Importantly, blocking visual experience during this developmental window by rearing mice 67 in complete darkness significantly impedes the maturation and eventual function of visual 68 circuits, whereas blockade of experience outside of this time frame does not have a strong 69 observable effect on circuit wiring<sup>11-13</sup>. Thus, sensory experience drives the 70 71 developmental refinement of thalamocortical circuits selectively during a defined window 72 of postnatal brain development in the mouse visual cortex.

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74 While the visual system has provided numerous essential insights into functional aspects of synaptic refinement and plasticity, our understanding of the molecular 75 mechanisms that mediate the influence of visual experience on circuit wiring remains 76 limited. One likely mechanism linking experience to circuit development in V1 is the 77 induction of gene programs in neurons in response to sensory-driven neuronal activity, a 78 79 process termed activity-dependent transcription<sup>14</sup>. In this process, synaptic innervation onto a given neuron drives the influx of Ca<sup>2+</sup> into the cell through ionotropic glutamate 80 receptors and L-type Ca<sup>2+</sup> channels<sup>15</sup>. This influx of Ca<sup>2+</sup> initiates intracellular signaling 81 cascades that phosphorylate, and thereby activate, transcription factors in the nucleus, 82 such as CREB and MEF2<sup>16,17</sup>. Within the first hour of neuronal activation, these factors 83 induce the expression of immediate early genes (IEGs), many of which encode a separate 84 85 set of transcription factors, including the well-established IEG Fos<sup>18</sup>. During a second wave of activity-dependent transcription which typically occurs between two and six hours 86 87 following neuronal activation, IEGs bind a subset of genomic promoters and enhancers 88 to drive the expression of a separate cohort of genes (late-response genes, LRGs) 89 encoding direct mediators of synaptic remodeling, such as the secreted neurotrophin Bdnf<sup>19,20</sup>. This two-wave pattern of activity-dependent transcription encompassing the 90 91 early expression of transcriptional regulators (i.e. IEGs) followed by the later expression of molecules that act locally at individual synapses (i.e. LRGs) is likely to contribute to the 92

refinement of visual circuits at the molecular level. Although a detailed analysis of 93 sensory-driven transcription during the critical window of circuit development in V1 has 94 not yet been established, work in adult animals suggests that the gene programs induced 95 by experience in this brain region are highly cell-type-specific, reflecting the capacity of 96 97 activity-dependent genes to shape cellular function in a precise manner and necessitating investigation at cell-type resolution<sup>21-23</sup>. Thus, genes that are induced by neurons in 98 response to sensory experience are promising candidates to mediate circuit development 99 100 during a period of sensory-dependent refinement in V1.

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102 Given the potential of sensory-induced gene programs to represent a key 103 molecular link between visual experience and circuit development, we reasoned that a 104 comprehensive atlas of sensory-dependent gene expression in developing V1 would provide useful insights into how circuits mature in response to environmental cues at the 105 106 molecular level. To this end, we leveraged a sensory deprivation and stimulation paradigm, the late-dark-rearing (LDR) paradigm, to dampen or increase visual experience 107 108 between P20 and P27, when sensory-dependent synapse remodeling begins and peaks. 109 respectively. We then performed single-nucleus RNA-sequencing (snRNAseq) on V1 110 tissue across six conditions encompassing normally reared, sensory deprived, and four 111 cohorts of sensory stimulated mice, followed by differential gene expression analysis to 112 identify transcripts that exhibited significantly greater RNA abundance in cells from stimulated versus unstimulated animals. We then further demonstrated the utility of the 113 dataset by comparing sensory-dependent gene programs between cell types, 114 115 characterizing transcriptional induction and repression events based upon RNA velocity, and predicting molecular interactions between cell types using the computational tool 116 *CellChat.* We expect this dataset, which describes sensory-driven changes in gene 117 118 expression across 118,529 nuclei representing sixteen distinct types of brain cells, to be 119 a valuable resource for investigators interested in uncovering the molecular basis of 120 sensory-dependent synapse remodeling and plasticity in the developing brain. 121

- 122 **Results**:
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A visual deprivation and stimulation paradigm for capturing sensory-induced transcripts 125

To characterize the gene programs that are elicited by sensory experience during 126 a critical period of postnatal brain development, we harnessed a dark-rearing method to 127 128 manipulate visual experience in mice in a temporally restricted manner. In this paradigm, 129 mice were initially reared according to a standard 12-hour light/12-hour dark cycle (normal 130 rearing, NR), the environment maintained in most animal facilities, before being placed in 131 complete darkness at P20, the beginning of sensory-dependent visual circuit refinement. Mice were then maintained in a completely dark environment for 24 hours a day across 132 133 a seven-day period. At P27, when sensory-dependent refinement peaks, one cohort of 134 dark-reared mice was sacrificed in the dark without re-exposure to light, and V1 tissue 135 was collected. Other cohorts of dark-reared mice were acutely re-exposed to white light at P27 for varying amounts of time following the week-long period of darkness, a 136 137 manipulation that leads to the acute and robust activation of circuit refinement and plasticity in both the dLGN and V1<sup>6,11,24</sup>. Thus, this late-dark-rearing (LDR) paradigm 138

139 allowed us to assess the impact of (1) sensory deprivation, and (2) sensory stimulation on gene expression selectively during a time when such experience is necessary for 140 neural circuit refinement (Fig. 1A). 141

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143 Previous studies of sensory-dependent transcription in the visual system have 144 focused on analyzing gene expression at two timepoints following the re-exposure of adult mice to light after dark-rearing (compared to unstimulated, sensory deprived mice): one 145 hour, when immediate-early genes (IEGs) have been proposed to peak, and four hours, 146 when late-response genes (LRGs) are highly induced<sup>21</sup>. However, given that these 147 studies were performed in adult mice, we first sought to confirm that these time points 148 149 were also optimal for capturing sensory-dependent gene expression during postnatal 150 development. To do so, we performed gPCR and single-molecule fluorescence in situ hybridization (smFISH) in parallel on V1 tissue after subjecting mice to the LDR paradigm 151 152 described above, and we assessed the expression of the canonical IEGs Fos and Jun as a read-out for the timing of sensory-dependent transcription. We found that the 153 expression of both IEGs was increased as early as fifteen minutes after light re-exposure 154 155 (i.e. acute sensory stimulation), and that this increase in Fos and Jun persisted for at least 156 two hours after stimulation. Within this time frame, the peak of Fos and Jun expression 157 occurred not at one hour but at thirty minutes after light re-exposure (Fig. 1B-D). Thus, 158 we included a thirty-minute stimulation timepoint to capture IEGs in our experiments. We also included three additional stimulation timepoints at which we expected to capture the 159 160 bulk of LRGs: two hours, four hours, and six hours. An additional benefit of including three late-response conditions is that it allowed us to derive insights into the dynamics of 161 sensory-dependent gene programs on a broader scale. Altogether, our finalized dataset 162 includes cells from mice according to the following six conditions: normally reared (NR) 163 mice at P27, mice reared in complete darkness from P20 to P27 (LDR), and mice reared 164 165 in complete darkness between P20 and P27 then acutely re-exposed to light for 30 166 minutes (LDR30m), two hours (LDR2h), four hours (LDR4h), or six hours (LDR6h). To our knowledge, this is the most extensive time course of sensory-dependent gene expression 167 168 at single-cell resolution generated to date.

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- 170 Mapping sensory-dependent gene expression in the developing cortex
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To map sensory-dependent changes in gene expression across cortical cell types 172 in an unbiased manner, we performed single-nucleus RNA-sequencing (snRNAseg: 10X 173 174 Genomics) on V1 tissue bilaterally micro-dissected from mice following the LDR paradigm 175 described above (Fig. 1A). We sequenced individual nuclei rather than whole cells based 176 upon our interest in capturing nascent transcriptional events that are acutely induced by 177 experience. Three biological replicates were performed for each condition, with each replicate being made up of cells pooled from the visual cortices of three animals to 178 179 increase yield. Biological replicates were collected, isolated, and processed 180 independently on different days to control for batch effects. After next-generation 181 sequencing, the data were mapped to the mouse genome and quality control was performed to remove putative doublets, unhealthy or dying cells, and droplets containing 182 183 ambient RNA from the dataset using Seurat, DoubletFinder, and DecontX packages in 184 R<sup>25-27</sup>. Data were then integrated across biological replicates and conditions for

downstream analysis within Seurat. The final dataset includes 118,529 nuclei across 16 185 distinct cell clusters representing eight excitatory neuron subtypes, four inhibitory neuron 186 subtypes, and four glial subtypes (Fig. 1E-G). The excitatory populations captured include 187 layer 4 (L4) pyramidal (PYR) neurons, layer 2/3 (L2/3) PYR neurons, three populations 188 189 of layer 5 (L5) neurons, and three populations of layer 6 (L6) neurons. Inhibitory 190 populations sequenced include Grin3a-enriched neurons (some of which also express SST markers), parvalbumin (PV) neurons, VIP neurons, and neurons expressing Npy, 191 which include neurogliaform cells (also positive for Lamp5) and a subset of SST neurons. 192 Glial populations sequenced include astrocytes, oligodendrocytes, oligodendrocyte 193 194 precursor cells (OPCs), and microglia (Fig. 1H,I). Cell type assignments were based upon 195 the presence of marker genes identified previously<sup>21,28</sup>. The numbers of cells within each 196 cell class included in the dataset are given in Table 1.

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#### 198 Sensory deprivation upregulates a cohort of genes in excitatory neurons

200 With this dataset in hand, we set out to understand how manipulating sensory 201 experience impacts the transcriptional states of cells in V1 during development. To this 202 end, we utilized the DEseg2 function within Seurat to identify transcripts that were 203 significantly differentially expressed (differentially expressed genes, DEGs; false 204 discovery rate (FDR) < 0.05) between each condition for each cell type, beginning with a comparison of gene expression in normally reared (NR) mice at P27 versus sensory 205 deprived (i.e. LDR) mice at the same age. This analysis revealed changes in gene 206 expression meeting a minimum threshold of  $log_2(1.5)$  fold change in excitatory neurons 207 following dark-rearing compared to NR mice. Specifically, when the DEG analysis was 208 209 applied to all excitatory neuron clusters in aggregate, 52 genes (e.g. the transcription 210 factor Stat4 and the cytoskeletal regulator Clmn) were found to be less highly expressed 211 in the NR condition compared to LDR mice, indicating that depriving mice of light 212 *increased* the expression of a define cohort of genes (Fig. 2A). Among excitatory neuron 213 clusters, the subtypes that exhibited the largest numbers of gene expression changes following LDR were L2/3 PYR neurons (86 genes upregulated after dark-rearing and 7 214 215 genes downregulated; Fig. 2B) followed by neurons in the L6a cluster (21 genes upregulated after dark-rearing; Fig. 2C). In L2/3 neurons, genes more highly expressed 216 217 in the LDR condition included factors such as Tspan11 and Gpc3 which are involved in cellular dynamics and migration<sup>29,30</sup>. Interestingly, the seven genes that were more highly 218 expressed in the NR condition included known activity-regulated factors such as the 219 220 neurotrophin Bdnf and the nuclear orphan receptor Nr4a1. In contrast to excitatory 221 neurons, only one gene, the synaptically localized long non-coding RNA Gm45323, was 222 less highly expressed in inhibitory neurons in the NR compared to the LDR condition<sup>31</sup> 223 (Fig. 2D). These findings suggest that excitatory neurons are more sensitive to the effects 224 of sensory deprivation than inhibitory neurons at the transcriptional level, and indicate 225 that sensory deprivation tends to increase, rather than decrease, gene expression in 226 excitatory neurons of V1.

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228 Excitatory and inhibitory neurons mount shared and distinct transcriptional responses to 229 sensory stimulation

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231 We next compared DEGs between each light re-exposure timepoint (LDR30m, 232 LDR2h, LDR4h, and LDR6h) and the sensory deprived LDR condition for all 16 cell 233 clusters in isolation, and for inhibitory and excitatory populations after aggregation. These 234 experiments revealed bidirectional changes in gene expression at every timepoint 235 analyzed within most cell types, yielding a total number of 1,268 unique genes that are 236 upregulated at any stimulation timepoint compared to the LDR condition (Table 2). These 237 genes included numerous previously identified activity-dependent IEGs, such as the AP1 238 factors Fos and Jun, the neuron-specific IEG Npas4, and the Nr4a and Eqr families of TFs that are induced by various extracellular stimuli including synaptic innervation. 239 240 Interestingly, although AP1 transcription factors are broadly considered to be IEGs, Jund 241 and Junb both exhibited a pattern of induction more consistent with an LRG identity, 242 peaking at LDR2h rather than LDR30m (Fig. 3A). Among all cell types analyzed, sensory 243 experience elicited the most robust gene expression changes in L2/3 and L4 excitatory 244 neurons, a result that we validated by performing smFISH for the IEGs Fos and Nr4a1 in 245 these layers (Fig. 3B-E). Both of these excitatory populations are innervated by projection 246 neurons outside of V1, with L2/3 neurons receiving top-down information from other 247 cortical areas and L4 neurons receiving bottom-up information from the visual thalamus 248 that is strongly driven by sensory experience. That L2/3 PYR neurons exhibit the largest 249 number of transcriptional changes as a result of light re-exposure is consistent with a 250 recent report identifying L2/3 cells as being particularly sensitive to sensory experience during postnatal development<sup>32</sup>. In addition to IEGs, we also identified cohorts of genes 251 252 that were preferentially upregulated at LDR2h, LDR4h, or LDR6h, fitting the expected profile of LRGs (Fig. 3F). Assessing changes in gene expression following light re-253 254 exposure in aggregated excitatory and inhibitory neuron populations revealed that the 255 most robust change in gene expression for excitatory neurons (233 genes upregulated) 256 occurred at the 30-minute timepoint, while the most robust change in inhibitory neurons 257 (82 genes upregulated) occurred six hours after light re-exposure (Fig. 3G). This 258 observation could reflect a temporal trajectory in which excitatory neurons are more 259 strongly impacted by sensory stimulation first, with inhibitory neurons responding later. 260

261 Studies in adult mice have suggested that IEGs tend to be conserved between cell 262 types while LRGs are more likely to be cell-type-specific. Thus, we next assessed the 263 overlap between the DEGs that were upregulated by stimulation at each time point across aggregated inhibitory and excitatory clusters. Unexpectedly, of the 233 genes that are 264 upregulated in excitatory neurons following light re-exposure at LDR30m, only 31 (13.3%) 265 266 were also upregulated in inhibitory cells at the same timepoint. Conversely, 47 (66%) of 267 the 71 genes upregulated in excitatory neurons at LDR6h were shared with inhibitory 268 neurons (Fig. 3H,I). These findings suggest one or both of two possibilities: (1) the LRG 269 programs within these cell types have more in common than earlier waves of sensory-270 induced transcription; and/or (2) inhibitory neurons may respond more slowly to visual 271 stimulation than excitatory neurons. The latter possibility is consistent with the finding that 272 excitatory neurons are more sensitive to sensory deprivation than inhibitory neurons at 273 the transcriptomic level (Fig. 2).

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The commonalities and distinctions between the gene programs induced by experience in excitatory versus inhibitory neurons were reflected in the functional 277 classifications of the sensory-induced genes identified in each cell type. For example, at 278 LDR30m, DEGs in both classes were enriched for gene ontology (GO) categories such 279 as RNA polymerase II-specific DNA-binding transcription factor binding, reflecting the 280 sensory-induced expression of members of the Nr4a family of nuclear orphan receptors 281 in both excitatory and inhibitory cells. Conversely, GO categories related to GTPase 282 binding and GTPase regulator activity, including the Rho GTPase guanine nucleotide exchange factors (RhoGEFs) Arhgef3 and Plekhg5, were selectively upregulated in 283 284 excitatory neurons at this timepoint (Fig. 3J,K). This observation suggests that excitatory 285 neurons may undergo structural remodeling as a result of sensory-dependent 286 transcription in a manner that is unique from inhibitory neurons, although these 287 transcriptional events would likely take longer than six hours to elicit a functional effect. 288 Overall, these data indicate that the gene programs induced in excitatory and inhibitory neurons downstream of sensory stimulation exhibit partial overlap at each time point 289 290 analyzed, with the amount and nature of overlap varying significantly by condition. These 291 observations support the utility of the snRNAseq dataset for uncovering transcripts that 292 are upregulated by sensory experience across multiple cell types as well as the 293 transcripts that are induced by experience in a cell-type-specific manner.

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295 Comparison of sensory-induced genes in L2/3 and L4 excitatory neurons reveals a 296 shared protein kinase signature and divergent axon guidance pathways

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298 Given that L2/3 and L4 neurons were among the most strongly impacted by 299 sensory experience, we next compared sensory-driven gene programs between these excitatory subpopulations. We first asked whether sensory-dependent gene programs in 300 301 L2/3 and L4 neurons share a general architecture by guantifying the numbers of genes 302 that were upregulated in both cell types following visual stimulation at each experimental 303 timepoint. For both cell classes, LDR30m was the time point at which the highest numbers 304 of genes (303 genes in L2/3 neurons and 239 genes in L4 neurons) were upregulated, 305 followed by LDR4h, when 210 and 124 genes were upregulated in L2/3 and L4 neurons, respectively (Fig. 4A-F). This observation suggested that, among the three later 306 307 timepoints analyzed, LDR4h was likely the peak of late-response, sensory-dependent transcription in L2/3 and L4 neurons. We next assessed the overlap between the gene 308 309 programs induced by L2/3 and L4 neurons at each timepoint. As expected, these gene programs exhibited substantial overlap. For example, of the combined 349 genes 310 upregulated at LDR30m across both cell classes, 193 (or 55%) were induced in both cell 311 312 types (Fig. 4G). Varying degrees of overlap were also observed between sensory-313 dependent gene programs at the later timepoints as follows: 31% overlap at LDR2h, 36% 314 at LDR4h, and 52% at LDR6h (Fig. 4H-J). Thus, L2/3 and L4 neurons mounted both 315 shared and distinct responses to sensory experience that were most robust at LDR30m 316 followed by LDR4h.

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We next investigated the nature of the sensory-dependent gene programs induced by both cell types by performing GO analysis on the sets of genes that were upregulated at LDR30m or LDR4h. As expected based upon the overlap in the gene programs induced in L2/3 and L4 neurons (Fig. 4G-J), several of the same GO categories emerged for both cell types. These shared functional classifications included *GTPase binding* (likely

reflecting mechanisms of cytoskeletal remodeling) and nuclear receptor binding 323 324 (associated with activity-dependent transcription factors), but the protein serine/threonine 325 kinase activity category was particularly prominently represented. In L2/3 neurons, this category was the most highly enriched functional classification of genes revealed by GO 326 327 analysis at LDR30m and the third most enriched at LDR4h (Fig. 4K,L). Similarly, protein 328 serine/threonine kinase activity is also the most enriched classification of upregulated 329 genes in L4 neurons at LDR30m (although not at LDR4h; Fig. M,N). Thus, genes encoding protein serine/threonine kinases were among the most strongly induced genes 330 following sensory stimulation in both L2/3 and L4 neurons, and they were induced as 331 332 early as LDR30m suggesting that their expression adheres to an IEG-like pattern. On the 333 contrary, these kinases were largely not induced by light re-exposure in inhibitory neurons 334 at LDR30m or LDR4h (Fig. 3K), suggesting that they may be particularly important for 335 mediating sensory-dependent plasticity in excitatory cells.

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337 While genes within the enriched protein serine/threonine kinase activity category 338 included those that encode intracellular molecules that regulate a wide range of cellular 339 processes, in neurons, these pathways are specialized to convey information about 340 changes at the cell membrane (and at synapses in particular) to the nucleus to shape gene expression following synaptic innervation<sup>33,34</sup>. For example, the Extracellular signal-341 342 regulated (ERK)-family kinases Mapk4 and Mapk6 were strongly upregulated by sensory stimulation at LDR30m in both L2/3 and L4 neurons, with Mapk4 representing one of the 343 most highly induced genes in L4 neurons at LDR4h. Likewise, the related Salt-inducible 344 kinases, Sik1-3, were among the most highly upregulated genes in both cell types at all 345 timepoints analyzed (Fig. 4C-F). Notably, intracellular signaling molecules, including ERK 346 347 and Sik family kinases, interact with numerous IEG transcription factors identified in the 348 dataset<sup>35-38</sup>. Thus, genes upregulated by sensory experience in L2/3 and L4 neurons 349 share a protein kinase signature that we predict may strengthen synapse-nucleus 350 crosstalk following sensory stimulation principally in excitatory neurons. 351

352 We next interrogated differences between the sensory-dependent gene programs 353 in L2/3 and L4 neurons by performing GO analysis on the gene sets that were uniquely 354 induced in each cell type. An interesting pattern to emerge was the differential induction 355 of two axon guidance pathways within these populations: the ephrin pathway (including 356 ephrin receptors *Ephb3* and *Epha10*) in L2/3 neurons and the semaphorin pathway 357 (including the semaphorin co-receptors *Plxna4* and *Nrp1*) in L4 neurons (Table 2). Both 358 of these pathways mediate the migration of neuronal axons and the establishment of 359 synapses within target zones based upon ephrin and semaphorin ligand expression, and have been implicated in establishing retinotopy in the developing visual system<sup>39-42</sup>. 360 361 These findings suggest that sensory experience may elicit axonal remodeling and/or presynaptic plasticity by inducing the expression of members of two distinct signaling 362 363 families, ephrins and semaphorins, in L2/3 and L4 neurons, respectively. This result is 364 consistent with the different projection patterns of these two cell types. 365

Sensory-dependent gene induction and repression dynamics in L2/3 and L4 neuronsrevealed by RNA velocity

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369 Increases in RNA abundance following sensory stimulation are often interpreted to 370 reflect the new transcription of genes. However, RNA abundance can be influenced by 371 many mechanisms beyond transcription, such as changes in the stability or degradation 372 of the mRNA. As single-cell transcriptomic approaches have evolved over the past 373 decade, new computational methods for disentangling transcriptional induction events 374 from other modes of gene regulation have become available. We applied one such 375 principle, RNA Velocity, to explicitly characterize genes whose expression is likely to be 376 upregulated by sensory stimulation via a transcriptional mechanism. Briefly, this approach estimates transient cell state transcriptional dynamics based upon the relative abundant 377 378 of nascent (unspliced) and mature (spliced) mRNAs.

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380 Given that the earliest versions of single-cell velocity (scVelo) analysis modules exhibited limited performance in identifying multiple rate kinetics (MURK) genes<sup>43,44</sup>. 381 382 which can exhibit rapid and complex changes at the transcriptional level, in our dataset, we applied a newer approach, UniTVelo, to assess RNA velocity in L2/3 and L4 neurons 383 across LDR and light re-exposure timepoints<sup>45</sup>. This framework utilizes a radial basis 384 385 function (RBF) model to overcome the limitations caused by the linear assumptions 386 employed by scVelo. UniTVelo's advanced approach allows for a more nuanced 387 understanding of gene expression patterns, accurately capturing the dynamics of genes 388 that display variable transcription rates at different stages of a cellular process. 389

390 Using UniTVelo, we first analyzed the architecture of transcript maturation for each predicted cell state transition: LDR to LDR30m, LDR30m to LDR2h, LDR2h to LDR4h, 391 and LDR4h to LDR6h. These comparisons revealed strong signatures of both 392 393 transcriptional induction and repression in L2/3 and L4 neurons with a stereotyped pattern 394 shared by both cell types (Fig. 5A). For example, between LDR and LDR30m, relatively large numbers of genes in each cell type (204 and 161 genes in L2/3 and L4 cells, 395 396 respectively) exhibited transcriptional induction with only very few genes exhibiting 397 repression at this timepoint. On the contrary, between LDR30m and LDR2h, the majority of significantly altered genes were repressed, not induced. Between LDR2h and LDR4h, 398 399 most altered genes were induced although many genes were also repressed. Finally, 400 between LDR4h and LDR6h, the majority of altered genes in each cell type exhibited a 401 repressed profile (Fig. 5B,C). These results are in line with the canonical view of stimulusdependent gene programs involving primarily two waves of transcription: an IEG wave 402 403 peaking at LDR30m and a LRG wave peaking at LDR4h.

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405 To more fully understand the dynamics underlying sensory-dependent 406 transcription, we next asked whether the genes that are induced at LDR30m exhibit 407 sustained expression across the timecourse or whether their expression returned to 408 normal by LDR6h. To do so, we compared the genes that were identified by UniTVelo as 409 induced between LDR and LDR30m to the genes that were repressed between LDR30m 410 and LDR2h for each cell type. Among the 204 genes that were induced in L2/3 neurons 411 between LDR and LDR30m, 112 (55%) were repressed between LDR30m and LDR2h 412 (Fig. 5D). A similar comparison in L4 neurons revealed that 38% of genes induced 413 between LDR and LDR30m are repressed between LDR30m and LDR2h (Fig. 5E). We 414 next compared the dynamics of genes that were upregulated at the LDR4h timepoint,

which our data suggests is the peak of LRG programs in both L2/3 and L4 neurons. We 415 observed that, among the 132 genes induced between LDR2h and LDR4h in L2/3 416 417 neurons, 98 genes (74%) were repressed between LDR4h and LDR6h (Fig. 5F). The 418 same analysis in L4 neurons revealed that, of the 45 genes that were induced between 419 LDR2h and LDR4h, 30 (67%) were repressed between LDR4h and LDR6h (Fig. 5G). 420 These data suggest that a significant proportion (between 38%-55%) of genes induced 421 at LDR30m are repressed relatively quickly within two hours of induction, while an even 422 more substantial proportion of the genes induced between LDR2h and LDR4h were 423 repressed at LDR6h. These data highlight distinct cohorts of genes in L2/3 and L4 424 neurons that exhibit transcriptional induction/repression dynamics within the time window 425 captured in our paradigm (Table 3).

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427 Given that the standard approach for defining sensory-induced genes is to 428 compare RNA abundance between a stimulated and an unstimulated condition (as 429 illustrated in Figs. 3 and 4), we next sought to determine what percentage of the genes 430 that were upregulated at LDR30m based upon total RNA abundance overlapped with the 431 genes identified as induced based upon RNA velocity. As expected, a greater number of 432 genes exhibited heightened expression at LDR30m versus LDR as determined by DEG 433 analysis in both L2/3 and L4 neurons than those identified by velocity as induced in either 434 cell type, suggesting that the list of DEGs for a given cell type in our dataset includes 435 genes that are upregulated via transcription-independent mechanisms, for example 436 possibly due to a decrease in the rate of that gene's mRNA degradation. However, 437 another interpretation could be that the DEG analysis is more sensitive than the RNA velocity approach. Nevertheless, we found that around 33% of the genes that were 438 439 upregulated between LDR and LDR30m in L2/3 and L4 neurons based upon DEG 440 analysis were also induced between these time points when assessed by RNA velocity 441 (Fig. 5H,I). A similar analysis of transcripts upregulated between LDR2h and LDR4h 442 revealed 27% overlap for L2/3 but only 11% for L4 neurons, suggesting that LRG programs in L4 neurons might be sustained longer than those in L2/3 neurons (Fig. 5J,K). 443 Consistent with overlap between the results of the DEG and velocity analyses. GO 444 445 analysis revealed that many of the same or similar ontological categories related to gene 446 function (e.g. nuclear receptor binding/activity and phosphatase binding/activity) were 447 found among the sets of genes upregulated in L2/3 neurons at LDR30m, with both categories also being among the most strongly enriched genes that are repressed 448 between LDR30m and LDR2h (Fig. 5L,M). A similar pattern emerged for L4 neurons, with 449 450 GO categories related to GTPase binding being induced then repressed (Fig. 4N,O). 451 Thus, the RNA velocity analysis uncovered subsets of DEGs that are most likely to 452 represent bona fide IEGs and LRGs based upon their transcriptional dynamics, exhibiting 453 the versatility of the snRNAseg dataset for understanding how sensory experience 454 modifies not just gene expression but transcription explicitly.

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Inference of cell-cell interactions using CellChat uncovers Neurexin and Neuregulinsignaling in developing V1

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459 Cells in the brain interact dynamically with one another not only through contact-460 mediated mechanisms but also through molecular signaling between compatible ligand-

receptor pairs. However, a systematic catalog of intercellular interactions in developing 461 visual cortex was lacking. Thus, we next harnessed our snRNAseq dataset to analyze 462 putative cell-cell interactions in V1 across all cell types using the computational tool 463 CellChat, which harnesses databases of known ligand-receptor binding partners to 464 465 estimate the number and strength of putative intercellular communication pathways based upon gene expression data<sup>46</sup>. Given our growing appreciation for the diversity of 466 brain cell types and their functions in circuit development, we applied CellChat to map 467 interactions between all cells in our dataset, focusing on the NR condition in which mice 468 are reared normally then processed at the peak of sensory-dependent refinement at P27. 469

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471 Applying CellChat to the NR condition within the snRNAseq dataset, we detected 472 442 significant ligand-receptor pairs among the 16 cell clusters captured (Fig. 6A,B). We 473 further categorized these pairs as belonging to 58 discrete signaling pathways. Consistent 474 with sensory experience promoting synaptic remodeling and maturation during the time 475 window analyzed, modules related to synapse development and plasticity were among 476 the strongest pathways identified. For example, the Neurexin (Nrxn) family of autism-477 linked presynaptic adhesion molecules that mediate synapse maintenance and function 478 by binding Neuroligins (NIgns) and Leucine-rich repeat transmembrane neuronal proteins 479 (Lrrtms) at postsynaptic specializations was the strongest signaling pathway uncovered 480 by CellChat. Signaling between Neurequlins (Nrgs) and ErbB receptors, which orchestrates the formation of excitatory synapses onto inhibitory neurons<sup>47,48</sup>, was the 481 second most enriched module identified. Apart from Nrxn and Nrg signaling, Ncam and 482 Cadm (i.e. SynCAM1) adhesion molecules were also identified as active signals in V1. 483 484 Furthermore, consistent with axonal remodeling occurring during sensory-dependent 485 refinement, EphA and EphB ephrin receptors and Semaphorins 3-6 were also predicted 486 to signal actively (Fig. 6C,D). These findings suggest that cells in V1 work together to 487 shape developing circuits in response to sensory experience via molecular signaling 488 pathways that converge upon synapses. 489

490 We next assessed the putative contributions of the different cell types in V1 to the Nrxn and Nrg signaling pathways identified via CellChat. The primary outgoing signals of 491 the Nrxn pathway were Nrxns 3 and 1, and they were most prominently expressed by L6b 492 493 excitatory neurons (Fig. 6C and 7A). The primary receivers of these signals were NIgn1 494 and Lrrtm4, which were most highly expressed in L5-PT neurons but also appeared in L4 495 neurons and to a lesser extent in other populations as well (Fig. 6D and 7A). In general, 496 we found that excitatory neurons were more heavily involved in both the propagation of 497 outgoing and the receipt of incoming molecular signals than interneurons or glia, with 498 neurons in L6 being particularly active in this regard (Fig. 7A,B). Interestingly, while the 499 inducible gene programs in L2/3 and L4 excitatory neurons shared many features (Fig. 500 4), these cell classes differed substantially in their participation in cell:cell signaling, with 501 L4 neurons being much more likely to participate in signaling with other V1 cells than L2/3 502 neurons. Among inhibitory populations, Npy-expressing cells were the strongest senders 503 of outgoing signals while VIP neurons were the strongest receivers (Fig. 7A,B). In contrast, several excitatory populations were predicted to produce Nrg with L6b neurons 504 505 being the most prominent expressers followed by L4 neurons. All inhibitory cells were 506 relatively strong receivers of Nrg signaling except for VIP neurons (Fig. 7B; Table 4).

507 Overall, these data highlight the utility of the snRNAseq resource described here to 508 uncover important principles underlying the molecular control of circuit maturation in the 509 developing brain.

510

## 511 **Discussion**:

512

513 Since the seminal work of Nobel laureates David Hubel and Torsten Wiesel in the 1960s<sup>49,50</sup>, sensory experience has been known to be a major driver of brain 514 development. However, our understanding of the molecular mechanisms engaged by 515 516 experience to shape brain wiring has remained limited. While molecular adaptations at 517 individual synapses, such as changes in neurotransmitter receptor composition, are well poised to mediate the effects of activity on a neuron's synapses within an acute time 518 519 frame, in a developmental context, more global adaptations are warranted. To this point, 520 the idea that robust changes in gene expression driven by sensory stimulation during brain development may play a vital role in circuit refinement is consistent with emerging 521 522 evidence that neurons in visual cortex undergo significant epigenetic and genomic 523 changes across the first month of life in mice, including between P20 and P27<sup>51,52</sup>. 524 Because these changes in gene expression occur at the cellular rather than the synaptic 525 level, they are likely to exert substantial influence over the development and maintenance 526 of circuits in the long-term.

527

528 Inspired by this idea, we here present a whole-transcriptome atlas of sensorydependent gene expression across 118,529 nuclei representing 16 distinct cell types in 529 the brain. We envision several ways in which this dataset can be used to increase our 530 531 understanding of brain development. For example, investigators interested in 532 understanding the role(s) of one or more specific genes in brain development can 533 determine whether their genes of interest are expressed in a sensory-dependent manner, 534 and, if so, which cell types upregulate their expression in response to experience. 535 Second, investigators can determine how specific cell types of interest modify their transcriptional profiles in response to sensory stimulation. Finally, given that the dataset 536 537 includes data from control mice reared normally until P27, investigators can use this data 538 to explore gene expression in V1 in the absence of manipulations of experience.

539

540 Several observations that we have made in interrogating this dataset may be of 541 particular interest for future studies. For example, the observation that L2/3 and L4 542 neurons strongly upregulate intracellular signaling molecules such as protein 543 serine/threonine kinases (including ERK and Sik family members) as early as 30 minutes 544 after stimulation suggests that sensory-dependent gene programs in these cells may 545 reinforce synapse-to-nucleus crosstalk, strengthening the ability of synaptic innervation 546 to shape the neuronal transcriptome. In addition, the observation that excitatory neurons 547 are likely more sensitive to experience than inhibitory cells, both at the level of sensory-548 induced gene expression changes as well as cell signaling interactions, could increase 549 our understanding of the differential roles that these cell types play in visual function. 550 Among excitatory neurons, the discovery that L2/3 neurons are particularly strongly 551 affected is consistent with a recent study highlighting that the maturation of these cells is influenced visual experience<sup>32</sup>. At the level of cell signaling, our data showing that the 552

553 strongest signatures were related to Nrxn and Nrg signaling pathways suggests that cellular interactions within developing V1 converge upon synapses. Altogether, we expect 554 this dataset and our experimental investigation thereof to serve as tools for investigators 555 556 interested in uncovering molecular mechanisms guiding sensory-dependent refinement 557 in the developing brain.

#### 558 **Methods** 559

- 560
- 561 Animals

562

All experiments were performed in compliance with protocols approved by the 563 Institutional Animal Care and Use Committee at Cold Spring Harbor Laboratory (CSHL). 564 Male C57BI/6J mice were obtained from the Jackson Laboratory (Cat #000664) then 565 566 housed at CSHL in an animal facility where average temperatures and humidity were 567 maintained between 68-70° Fahrenheit and 54-58%, respectively. Mice in this study were aged between P18 – P27. Animals had access to food and water ad libitum. 568

- 569
- 570 Late-dark-rearing (LDR) paradigm

571 572 Male C57BI/6J mice were obtained from the Jackson laboratory at P18 and 573 allowed to acclimate to the standard 12-hour light/12-hour dark environment of the CSHL 574 animal facility until P20, at which point they were separated into six cohorts. One cohort 575 was maintained under normal housing conditions (normally reared, NR) while the other 576 cohorts were placed inside a well-ventilated, 100% light-proof chamber (Actimetrics). Mice 577 in the chamber were housed in complete darkness until P27, at which point one cohort 578 was sacrificed and perfused with ice cold 1X PBS (snRNAseg experiments) or 1X PBS 579 followed by 4% PFA (smFISH experiments) in the dark. The remaining four cohorts of 580 mice were also dark-reared between P20 and P27 but were then re-exposed to light for 581 varying lengths of time: 30 minutes (LDR30m), two hours (LDR2h), four hours (LDR4h), and six hours (LDR6h). After perfusing the mice and removing their brains in the dark, V1 582 regions were micro-dissected from all cohorts in the wet lab. 583

584

#### 585 Single-nucleus RNA sequencing and data analysis

586 V1 tissue collection

587 Whole brains were placed into ice-cold 1X Hank's Balanced Salt Solution (HBSS) supplemented with Mg<sup>2+</sup> and Ca<sup>2+</sup>. The V1 brain regions were then bilaterally micro-588 dissected under a 3.5X-90X Stereo Zoom microscope (AmScope) using a needle blade. 589 590 Micro-dissected tissue was either immediately processed for snRNAseg or was frozen for 591 later processing.

- 592 Nuclear suspension preparation

593 594 The V1 tissue was transferred to a 1 mL dounce homogenizer containing 300 µL

595 of ice-cold supplemented Homogenization Buffer (0.25M Sucrose, 25mM KCl, 5mM 596 MgCl<sub>2</sub>, 20mM Tricine-KOH, 5mM DTT, 0.75mM Spermine, 2.5mM Spermidine, 0.05X

597 Protease Inhibitor Cocktail,  $1U/\mu L$  of Rnase Inhibitor and 0.15% IGEPAL CA-630). Note 598 the inclusion of drugs to block gene transcription and protease activity, as well as an 599 RNase inhibitory to protect the integrity of the RNA. The tissue was homogenized with a 600 loose and tight pestle about 10-15 times, respectively. The samples were then filtered 601 using a 20  $\mu$ m filter.

## 602 Library construction and sequencing

603
604 Single-cell gene expression libraries were prepared using the Single Cell 3' Gene
605 Expression kit v3.1 (10× Genomics, #1000268) according to manufacturer's instructions.
606 Libraries were sequenced on an Illumina Nextseq2000 to a mean depth of ~30,000 reads
607 per cell.

608

#### 609 *Raw data processing*

The raw FASTQ files were processed using Cell Ranger (v7.1.0) and aligned to the mm10 reference mouse genome. Loom files for cell dynamics analysis were generated using Velocyto (v0.17.17) by mapping BAM files to the gene annotation GTF file (refdata-gex-mm10-2020-A). Each library derived from the single-nucleus datasets underwent identical processing, resulting in a gene expression matrix of mRNA counts across genes and individual nuclei. Each cell was annotated with the sample name for subsequent batch correction and meta-analysis.

#### 617 Quality control, cell clustering, and cell type annotation

To ensure the integrity of our single-cell RNA sequencing data, we implemented 618 619 several quality control measures. First, we calculated the log10 of the number of genes 620 per UMI (log10GenesPerUMI), and cells with a value less than 0.85 were excluded. We 621 also removed cells with more than 1% mitochondrial gene expression to reduce noise 622 from apoptotic or damaged cells. Additional thresholds included excluding cells with fewer than 500 UMIs or 300 genes to eliminate low-quality or empty droplets. Doublets were 623 624 identified and excluded using the DoubletFinder package, with optimal pK values determined for each sample through a sweep analysis<sup>25</sup>. Ambient RNA was removed with 625 DecontX<sup>27</sup>. Following these steps, we applied the standard Seurat (v4) pipeline for data 626 627 pre-processing (https://satijalab.org/seurat/articles/get started.html), which included 628 selecting the top 3,000 highly variable genes and regressing out UMI counts and mitochondrial gene percentage for cell clustering. 629

630 Clustering utilized the functions *FindNeighbors* and *FindClusters* from Seurat, 631 employing resolutions ranging from 0.1 to 0.5. A resolution of 0.5 was ultimately selected 632 for clustering. To identify major cell types, the *ConserveredMarkers* function (log2 fold 633 change > 0.25, MAST test, adjusted p-value < 0.05 with Bonferroni correction), with 634 pct.1 > 70% and pct.2 < 30% identified unique and highly enriched differentially expressed 635 genes (DEGs) in specific clusters compared to others. Cell types were manually

annotated based on the expression of conserved markers<sup>21,28</sup>, ensuring precise
 identification and accurate analysis of cellular phenotypes.

638 DEG analysis

Differentially expressed genes (DEGs) between conditions were identified using
 the DEseq2 function within Seurat v4. DEGs were identified for each of the 16 individual
 clusters included in the dataset.

642 RNA velocity analysis

643 Cell velocity analysis was conducted on L2/3 and L4 excitatory neurons using the 644 UniTVelo (v0.2.4) tool within the scVelo (v0.2.5.) framework, focusing on the 2000 most 645 variably expressed genes. Genes were categorized based on their fit\_t scores such that 646 those with a fit\_t>0 were classified as induced genes, whereas genes with a fit\_t <0 were 647 identified as repressed genes.

648 Cell-cell interaction analysis

The R package *CellChat* (http://www.cellchat.org/) was utilized to infer cell-cell interactions within our dataset. We adhered to the standard pipeline and default parameters set by *CellChat*. The complete CellChatDB.mouse database was employed, which categorizes ligand-receptor pairs into "Secreted Signaling," "ECM-Receptor," and "Cell-Cell Contact." Additionally, we conducted CellChat analyses on the overall dataset and separately for conditions at specific timepoints—LDR0, LDR30m, LDR2h, LDR4h, LDR6h, and NR, although we focus on the NR condition in this paper.

656 Enrichment analysis

657 Gene Ontology (GO) enrichment analysis was conducted using the "clusterProfiler"(v4.10.0) package. For the analysis of differentially expressed genes 658 (DEGs), only genes with an adjusted p-value less than 0.05 and a log2 fold change 659 660 greater than log2(1.5) were included. For the analysis of induced and repressed genes, 661 all identified genes were considered. The parameters for the GO analysis were set with a p-value cutoff of 0.05 and a q-value cutoff of 0.2, using the Benjamini-Hochberg (BH) 662 663 method for adjusting p-values. This approach ensures rigorous identification of biological 664 processes significantly associated with the gene sets under study.

- 665 *Real-time qPCR*
- 666

Flash frozen V1 samples were processed for RNA extraction using Trizol (ThermoFisher cat #15596018) according to the manufacturer's protocol. The cDNA library was built using iScript Kit (BioRad, cat #1725037) and Oligo d(T) primers. The realtime PCR were performed using SybrGreen kit (Fisher, cat #A25742) and standard PCR temperature protocol. *Fos* and *Jun* expression were normalized to *Gapdh* levels. The following primer sequences were used.

673

- 674 *Fos* (Forward): 5'-GGGAATGGTGAAGACCGTGTCA-3'
- 675 *Fos* (Reverse): 5'-GCAGCCATCTTATTCCGTTCCC-3'
- 676 Jun (Forward): 5'-CAGTCCAGCAATGGGCACATCA-3'
- 677 Jun (Reverse): 5'-GGAAGCGTGTTCTGGCTATGCA-3'
- 678 *Gapdh* (Forward): 5'-CATCACTGCCACCCAGAAGACTG-3'
- 679 *Gapdh* (Reverse): 5'-ATGCCAGTGAGCTTCCCGTTCAG-3'680
- 681 Single-molecule fluorescence in situ hybridization (smFISH)
- 682

683 Animals were anesthetized with a ketamine and xylazine cocktail (K: 90 mg/kg X: 684 10 mg/kg) before perfusion with ice-cold phosphate-buffered saline (PBS) followed by 4% paraformaldehyde (PFA) in 1X PBS. Brains were then drop-fixed in 4% PFA in 1X PBS 685 for 24 hours. Brains were then washed with 1X PBS thrice for 10 min before being 686 transferred to a 30% sucrose solution at 4° C. After dehydration, brains were embedded 687 688 in optimal cutting temperature (OCT; VWR cat #25608-930) and stored at -80° C. 20-µm 689 thick coronal sections containing the visual cortex were cut using a cryostat and thaw-690 mounted onto a Superfrost Plus microscope slide (Thermo Fisher Scientific, cat #1255015) and stored at -80° C until the experiment. FISH was performed using the 691 692 RNAScope platform V2 kit (Advanced Cell Diagnostics (ACD), cat #323100) according to 693 the manufacturer's protocol for fixed-frozen sections. Samples were then counterstained 694 with DAPI before ProLong Gold Antifade was applied. A 1.5X thickness coverslip was 695 then applied to the slides which were then stored at 4° C until imaging. Commercial 696 probes from ACDBio were obtained to detect the following genes: Fos (316921), Nr4a1 697 (423342-C2), and Jun (453561-C3).

- 698
- 699 *Confocal Imaging* 700

smFISH images were acquired using the Zeiss LSM780 with a x20/0.8 objective.
 Z-stack images were acquired.

- 703
- 704 FISH Quantification 705

FISH images were analyzed using FIJI. For each image, ROIs of layer 4 and layer 2/3 of the visual cortex were defined. The mean gray values were then taken for each ROI. For each mouse, the average mean gray value across both hemispheres was analyzed for both layer 4 and layer 2/3. A 2-way ANOVA was performed to test for significance.

- 711
- 712 Data availability
- 713

Both raw and processed snRNA-seq data are available at Gene Expression Omnibusunder accession number GSE269482.

- 716
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728 Declaration of interests

729

The authors declare no conflicts of interest.

# 732 Author contribution statement

733

731

AMX and LC conceptualized the study. AMX and CK designed, optimized, and performed snRNAseq. AMX performed and analyzed qPCR. CK performed and analyzed fluorescence *in situ* hybridization. QL and AMX analyzed the snRNAseq data, and QL and LC generated figures. LC wrote the paper.

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# 883 Figure Legends

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Figure 1. Experimental design and introduction to the single-nucleus RNA-885 886 sequencing dataset. (A) Schematic describing the late-dark-rearing (LDR) paradigm and the workflow of the single-nucleus RNA-sequencing (snRNAseg) experiments. (B) 887 Quantification of Fos mRNA expression in sensory deprived (LDR) mice and in mice 888 889 acutely exposed to light for between 15 minutes and 2 hours, with stimulation timepoints 890 labeled as follows: LDR15m (15 min of light), LDR30m (30 min), LDR1h (1 hour), and 891 LDR2h (2 hours). Fos expression assessed by qPCR and normalized to Gapdh 892 expression. Values plotted are additionally normalized to the LDR condition. (C) qPCR quantification of Jun mRNA expression (normalized to Gapdh) in V1 across all timepoints. 893 894 Data obtained by gPCR and values plotted are normalized to LDR. (D) Example confocal 895 images of V1 in sections from a sensory deprived mouse (LDR) and a mouse re-exposed 896 to light for thirty minutes (LDR30m). Fos mRNA (red), Jun mRNA (green), and DAPI (blue). Scale bar, 44 µm. (E) UMAP plot illustrating the 118,529 nuclei in the dataset 897 categorized by general cell class: excitatory neurons (periwinkle), inhibitory neurons 898 899 (salmon), and glia (green). (F) UMAP plot with all 16 clusters colored and labeled by cell

type. (G) UMAP plot with cells colored by condition according to the legend on the left. 900 901 Note that cluster composition is largely unaffected by sensory deprivation or stimulation. (H) Numbers of cells of each type included in the final dataset across all conditions. See 902 903 also Table 1. (I) Violin plot demonstrating the enrichment of markers used to assign nuclei 904 in the dataset to distinct cell types. Top enriched gene per cluster given on the Y-axis on 905 the right, normalized FPKM expression given on the Y-axis on the left, and cluster identity shown on the X-axis. For (B) and (C), n = 3 mice per condition; One-way ANOVA followed 906 by Tukey's post hoc test: \*p < 0.05; \*\*p < 0.01; \*\*\*p < 0.001; \*\*\*\*p < 0.0001. 907

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909 Figure 2. Sensory deprivation upregulates a cohort of genes in excitatory neurons.

910 (A) Volcano plot demonstrating transcripts that were significantly differentially expressed 911 (differentially expressed genes, DEGs) in aggregated excitatory neuron clusters after LDR compared to normally reared (NR) control mice. Y-axis, negative Log(10) adjusted 912 913 p value (threshold of p.adj < 0.05 indicated by dashed horizontal line). X-axis, Log(2) fold change (threshold of  $log_2(1.5)$  indicated by dashed vertical lines). Red, genes that are 914 915 more highly expressed in the NR condition (up in NR). Blue, genes that are more highly 916 expressed in the LDR condition (up in LDR). (B) Volcano plot of DEGs altered by sensory 917 deprivation in excitatory L2/3 neurons. (C) Volcano plot of DEGs altered by sensory 918 deprivation in L6a neurons. (D) Volcano plot of DEGs altered by sensory deprivation in 919 aggregated inhibitory clusters.

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921 Figure 3. Excitatory and inhibitory neurons mount shared and distinct responses to sensory stimulation. (A) Bubble plot illustrating the induction of canonical immediate-922 early genes (IEGs) across timepoints and cell types. Color indicates relative expression 923 924 level according to the scale on the right. Size of circle represents the percentage of cells 925 expressing the gene. (B) Confocal images of V1 in late-dark-reared (LDR) mice and in 926 mice re-exposed to light for 30 min (LDR30m) subjected to single molecule fluorescence in situ hybridization (smFISH) to label Fos mRNA. Scale bar, 100 µm. (C) Quantification 927 928 of Fos expression (arbitrary units, A.U.) in L2/3 and L4 of V1 in LDR and LDR30m mice. Two-Way ANOVA with Tukey's post hoc test. \*\*p<0.01, \*\*\*p<0.001; n = 3 mice/condition. 929 930 (D) Confocal images of V1 in LDR and LDR30m mice subjected to smFISH to label Nr4a1 mRNA. Scale bar, 100 µm. (E) Quantification of Nr4a1 expression in L2/3 and L4 in LDR 931 and LDR30m mice. Two-Way ANOVA with Tukey's post hoc test. \*\*p<0.01, \*\*\*p<0.001; n 932 933 = 3 mice/condition. (F) Bubble plot demonstrating late-response gene (LRG) expression 934 across cell types and conditions. Scaled expression indicated on the right. (G) Graph displaying the numbers of genes significantly upregulated at each stimulation timepoint 935 936 (compared to LDR control) across conditions for aggregated excitatory (salmon) and 937 inhibitory (periwinkle) neurons. (H) Venn diagram demonstrating overlap between 938 sensory-dependent gene programs in excitatory (salmon) versus inhibitory (periwinkle) neurons at LDR30m. (I) Venn diagram demonstrating overlap between sensory-939 940 dependent gene programs in inhibitory versus excitatory neurons at LDR6h. (J) Gene 941 ontology (GO) categories enriched among genes upregulated in excitatory neurons at 942 LDR30m. (K) GO categories enriched among genes upregulated in inhibitory neurons at 943 LDR30m. 944

Figure 4. Comparison of sensory-driven gene expression in L2/3 and L4 excitatory 945 946 neurons reveals a shared protein kinase signature and divergent axon guidance 947 pathways. (A) Schematic of the pathway from the retina to primary visual cortex (V1) in 948 the mouse. L2/3 neurons principally receive 'top-down' input from other regions of cortex 949 (blue), while L4 neurons receive "bottom-up" inputs from visual thalamus (magenta). (B) 950 Graph displaying the numbers of genes significantly upregulated at each stimulation timepoint (compared to late-dark-reared [LDR] control) across conditions for L2/3 951 (salmon) and L4 (periwinkle) neurons. (C) Volcano plot illustrating genes that were 952 953 significantly upregulated (red) or downregulated (blue) in L2/3 neurons after 30 minutes 954 of light re-exposure following LDR. (D) Volcano plot illustrating genes that were 955 significantly upregulated (red) or downregulated (blue) in L4 neurons after 30 minutes of 956 light re-exposure following LDR. (E) Volcano plot illustrating genes that were significantly 957 upregulated (red) or downregulated (blue) in L2/3 neurons after 4 hours of light re-958 exposure following LDR. (F) Volcano plot illustrating genes that were significantly upregulated (red) or downregulated (blue) in L4 neurons after 4 hours of light re-exposure 959 960 following LDR. (G) Venn diagram displaying overlap between upregulated genes 961 identified in L2/3 (salmon) versus L4 neurons (periwinkle) at the LDR30m timepoint. (H) 962 Venn diagram displaying overlap between upregulated genes in L2/3 versus L4 neurons 963 at the LDR2h timepoint. (I) Venn diagram displaying overlap between upregulated genes 964 identified in L2/3 versus L4 neurons at the LDR4h timepoint. (J) Venn diagram displaying overlap between upregulated genes identified in L2/3 versus L4 neurons at the LDR6h 965 966 timepoint. (K) Gene ontology (GO) analysis of genes upregulated by light in L2/3 neurons at LDR30m. (L) GO analysis of genes upregulated by light in L2/3 neurons at LDR4h. (M) 967 GO analysis of genes upregulated by light in L4 neurons at LDR30m. (N) GO analysis of 968 969 genes upregulated by light in L4 neurons at LDR4h. 970

971 Figure 5. Transcriptional induction and repression events in L2/3 and L4 neurons 972 revealed by RNA Velocity. (A) UMAP plots generated based upon RNA Velocity displaying transcriptional dynamics across each cell-state transition. L2/3 neurons, top 973 974 row, L4 neurons, bottom row. (B) Bar graph displaying the total numbers of induced (red) 975 and repressed (blue) genes across each cell-state transition in L2/3 neurons. (C) Bar graph displaying the total numbers of induced (red) and repressed (blue) genes across 976 977 each cell-state transition in L4 neurons. (D) Venn diagram displaying overlap between the 978 genes induced at LDR30m (red) and the genes that are repressed between LDR30m and 979 LDR2h (blue) in L2/3 neurons. (E) Venn diagram displaying overlap between the genes 980 induced at LDR30m (red) and the genes that are repressed between LDR30m and LDR2h 981 (blue) in L4 neurons. (F) Venn diagram displaying overlap between the genes induced 982 between LDR2h and LDR4h (red) and the genes that are repressed between LDR4h and 983 LDR6h (blue) in L2/3 neurons. (G) Venn diagram displaying overlap between the genes 984 induced between LDR2h and LDR4h (red) and the genes that are repressed between 985 LDR4h and LDR6h (blue) in L4 neurons. (H) Overlap between upregulated DEGs and 986 induced genes in L2/3 neurons at LDR30m. (I) Overlap between upregulated DEGs and 987 induced genes in L4 neurons at LDR30m. (J) Overlap between upregulated DEGs and induced genes in L2/3 neurons at LDR4h. (K) Overlap between upregulated DEGs and 988 989 induced genes in L4 neurons at LDR4h. (L) Gene ontology (GO) analysis of genes 990 induced in L2/3 neurons between LDR and LDR30m. (M) GO analysis of genes repressed

in L2/3 neurons between LDR30m and LDR2h. (N) GO analysis of genes induced in L4
 neurons between LDR and LDR30m. (O) GO analysis of genes repressed in L4 neurons
 between LDR30m and LDR2h.

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995 Figure 6. Inference of putative cell:cell interactions in developing V1 using 996 **CellChat.** (A) Cellular communication plot demonstrating the predicted numbers of 997 intercellular ligand-receptor interactions between all cell types in the dataset. (B) 998 Comparative weights/strength of the predicted cell:cell interactions plotted in (A). (C),(D) 999 Heatmaps displaying distinct cell signaling modules (y axis, pathways of interest in bold) 1000 predicted by *CellChat* across all cell types (x axis) in the dataset. Top, bars representing 1001 the contributions of each cell type to outgoing (C) or incoming (D) signals aggregated 1002 across signaling modules. Bar graphs on the right of each heatmap demonstrate the contribution of each individual signaling pathway to the overall interaction score 1003 1004 generated in *CellChat*. Heatmap colors indicate the relative strength of a given pathway's 1005 signaling activity as predicted by *CellChat* according to the scale on the right.

**Figure 7. Excitatory-excitatory signaling and excitatory-inhibitory signaling mediated by neurexin and neuregulin pathways, respectively**. (A) Hierarchical plot showing Nrxn-mediated interactions from excitatory to excitatory neurons (left) and from excitatory to inhibitory and glial cells (right). (B) Hierarchical plot showing Nrg-mediated interactions from excitatory neurons (left) and from excitatory to inhibitory and glial cells (right).

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1015 <u>Tables</u>: 1016

## 1017 **Table 1. Number of each cell type represented in the dataset.**

1018 Table 1 displays the number of cells included in the final dataset by condition and cell 1019 type. Cell types listed in alphabetical order from top to bottom.

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# 1021 Table 2. Differentially expressed genes identified in each cell type.

Table 2 includes all significantly differentially expressed genes identified by DEseq2 across cell types and conditions. Cell types and conditions listed in alphabetical order from top to bottom.

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# 1026 **Table 3. Induced and repressed genes based upon RNA velocity.**

- 1027 Table 3 includes genes identified as transcriptionally induced or repressed in L2/3 and L4 1028 excitatory neurons based upon RNA velocity.
- 1030 Table 4. Cell signaling modules in developing V1 identified by *CellChat*.
- 1031 Table 4 includes a list of ligand-receptor pairs identified via *CellChat*.
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Figure 1. Experimental design and introduction to the single-nucleus RNA-sequencing dataset.



Figure 2. Sensory deprivation upregulates a cohort of genes in excitatory neurons.

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Figure 3. Excitatory and inhibitory neurons mount shared and distinct responses to sensory stimulation.



Figure 4. Comparison of sensory-driven gene expression in L2/3 and L4 excitatory neurons reveals a shared protein kinase signature and divergent axon guidance pathways



Figure 5. Transcriptional induction and repression events in L2/3 and L4 neurons revealed by RNA velocity.



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