

Ire1 α -Regulated Rate of mRNA Translation is Required for Acquisition of Identity and Polarity in Upper Layer Cortical Neurons

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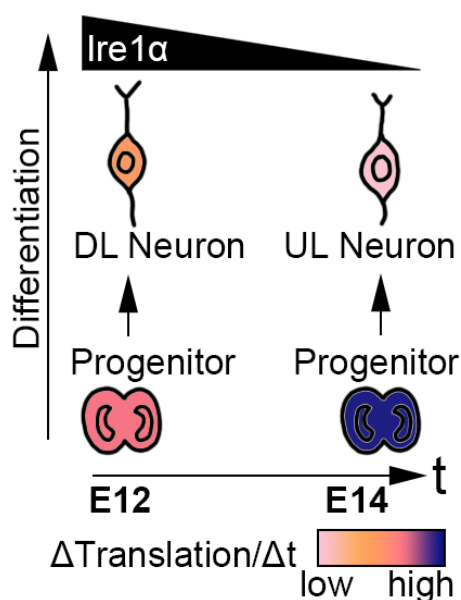
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SUMMARY Establishment of cortical layers and axon-dendrite polarity in neurons is fundamental for brain connectivity. Here, we present that timed mRNA translation control by Inositol-Requiring Enzyme 1 α , Ire1 α , is necessary for acquisition of upper layer neuronal identity and a single axon. We demonstrate that Ire1 α acts as a canonical regulator of global protein synthesis in developing cortical neurons, controlling the level of actively translating ribosomes, expression of translation factors, and ribosomal proteins. Translation rates distinguish early and late neuronal progenitors and early- and late-born postmitotic neurons, indicative of developmental stage- and differentiation-specific requirements for protein synthesis rates in the formation of upper and deeper cortical layers. We demonstrate that specification and polarization of upper layer neurons is uniquely sensitive to translation rate, in contrast to deep layer neurons. Our data shed light onto the post-transcriptional source of cellular diversity in the developing cortex and unveils stress-independent homeostatic functions of Ire1 α .

Ire1 α -Regulated Rate of mRNA Translation is Required for Acquisition of Identity and Polarity in Upper Layer Cortical Neurons



eTOC

Ambrozkiewicz and Borisova et al. demonstrate that upper layer identity and axon-dendrite polarity establishment requires translation rate regulation downstream of Ire1 α in the developing cortex.

HIGHLIGHTS

- Small molecule screening reveals Ire1 α upstream of upper layer neuronal identity
- Axon acquisition in cortical neurons requires Ire1 α
- Ire1 α -regulated translation rate is a feature of early and late cortical progenitors
- Transient block of translation inhibits axon formation and upper layer identity

KEYWORDS: cortical development, neuronal fate, upper layers, polarity, axon, protein translation, neuronal progenitors, Ire1 α , Satb2

INTRODUCTION

The provenance of cellular diversity in the cerebral cortex has been the center of attention for developmental neuroscientists for decades. The mammalian neocortex, evolutionarily the newest add-on to the central nervous system, is organized in six laminae, each populated by 5 neurons with distinct afferents and efferent connectivity patterns, transcriptomic signatures, and electrophysiological properties. Pyramidal glutamatergic neurons of the neocortex are born from the progenitors at the ventricular side of the brain primordium and polarize synchronously with their migration to laminate distinct cortical layers. Deeper layers, forming subcortical tracts, are established first. Later born, upper layer neurons migrate through pre- 10 established deeper layers, and differentiate to extend polarized axons forming intracortical tracts, such as the corpus callosum (Arimura and Kaibuchi, 2005; Barnes et al., 2007; Götz et al., 1998; Malatesta et al., 2000; Rakic, 1972; Tarabykin et al., 2001).

Transcription factor signatures characterize pyramidal neurons of different layers. Special AT-Rich Sequence-Binding Protein 2 (Satb2) is one of the strongest determinant of 15 intracortically projecting pyramidal neurons identity and is required to form the corpus callosum (Alcamo et al., 2008; Britanova et al., 2008). Conversely, subcortical projections are determined by expression of FEZ Family Zinc Finger (Fezf) and COUP-TF-Interacting Protein 2 (CTIP2, also known as Bcl11b) in deep layer V and VI neurons, where loss of either gene disrupts the corticospinal tract (Arlotta et al., 2005; Chen et al., 2005; McKenna et al., 2015; 20 Molyneaux et al., 2005).

The temporal progression of ventricular progenitor state requires restriction of multipotency towards generating solely upper layer Satb2-positive neurons. Current models describe the temporal progression of neuronal differentiation as conserved across deep and upper layers, but the initial ground states embedded in their progenitors as progressively 25 discrete (La Manno et al., 2020; Nowakowski et al., 2017; Telley et al., 2019). Transcriptomic signatures in apical progenitors are temporally plastic, and able to adapt to the developmental milieu in elegant heterochronic transplantation experiments (Oberst et al., 2019). Transcriptional priming in progenitors involves translational repression which restricts expression of early lineage genes in the later born lineage (Zahr et al., 2018).

30 Intrinsic genetic programs along with extrinsic cues further establish axon-dendrite polarity and cortical neuron connectivity. Single axons guided through postmitotic differentiation programs specify their postsynaptic targets (Barnes and Polleux, 2009). Current challenges include determining how intra- and extracellular determinants of polarity interplay with the neuronal fate programs to establish neuronal networks during development.

35 The models of neuronal fate and connectivity specification summarized above have focused heavily on transcriptional measurements, leaving the post-transcription as an understudied layer of regulation in neocortical gene expression. We have previously demonstrated that RNA-binding proteins, translation, and ubiquitination pathways are major regulators of cortical connectivity, neuronal subtype specification, and axonal projections 40 (Ambrozkiwicz et al., 2018; Kraushar et al., 2014). In the mature brain, high-fidelity protein translation is proven essential to maintain synaptic transmission (Biever et al., 2020; Hafner et al., 2019). However, how translation regulates key developmental events, including neuronal polarization and the temporal succession of cortical progenitors, remains unclear.

In this study, we sought upstream regulators of *Satb2*, which determines the
45 specification of callosally projecting cortical neurons (Alcamo et al., 2008; Britanova et al.,
2008), and represents the late-stage restricted output of multipotent progenitors. Using a
genetic reporter and Fluorescence Activated Cell Sorting (FACS), we screened a library of
small molecules for their ability to alter *Satb2* expression in developing neurons. Among our
most impactful candidates was an inhibitor of kinase/RNase Inositol-Requiring Enzyme 1 α
50 (*Ire1 α*), also known as ER-to-Nucleus Signaling 1 (*Ern1*), the main sensor of ER lumen
homeostasis and regulator of the Unfolded Protein Response (UPR). Upon ER stress, *Ire1 α*
signaling results in a translational shift promoting expression of proteins vital for cell survival
and restoration of the ER folding capacity (Yoshida et al., 2001). *Ire1 α* regulates stress-
independent remodeling of actin filaments by its association with filamin A (Urrea et al., 2018),
55 linked to periventricular heterotopia (Sheen, 2001). Another study unveiled a direct interaction
of *Ire1 α* with the ribosomes *in vitro* (Acosta-Alvear et al., 2018). Beyond that, homeostatic
functions of *Ire1 α* have remained quite elusive.

In this work, we demonstrate that *Ire1 α* is essential for acquisition of *Satb2* neuronal
identity during corticogenesis. In upper layer neurons, loss of *Ire1 α* is associated with loss of
60 axon polarity. Conditional deletion of *Ire1 α* in the developing neocortex changes the synthesis
of ribosomal proteins and the levels of actively translating ribosomes. Using fluorescent
noncanonical amino acid tagging (FUNCAT), we show that *Ire1 α* modulates translation rates
in cortical progenitors and neurons. Moreover, we reveal that the rate of protein synthesis
distinguishes early and late neuronal progenitors and their derived postmitotic neurons.
65 Transient inhibition of translation early during development results in loss of *Satb2* expression
and axon polarity. Taken together, our study defines translational signatures of neuronal
progenitors and distinct neuronal subtypes, unveiling the layers of gene expression regulation
driving neuronal diversity during corticogenesis. Additionally, this study extends the function
of *Ire1 α* beyond stress pathways to an innate developmental signal for neuronal specification
70 in the cortex.

RESULTS

***Ire1 α* is a positive regulator of *Satb2* and axon specification in developing cortical neurons**

75 In order to identify signaling pathways orchestrating neuronal subtype specification, we have
previously developed a method for screening small molecule inhibitors for their role in the
control of cell fate acquisition (Ambrozkiewicz et al., 2017), which utilizes the *Satb2*^{Cre/+}
mouse line (Britanova et al., 2006). By expressing a Cre-inducible fluorescent reporter loxP-
Stop-loxP-tdTomato in *Satb2*^{Cre/+} primary cortical cells (Fig. 1a), we used cell sorting to
80 quantify the proportion of *Satb2*-expressing tdTomato-positive cells (*Satb2*^{tdTom}) to EGFP-
expressing cells. We used EGFP-encoding reference plasmid to normalize for the
transfection efficiency.

We prepared *Satb2*^{Cre/+} primary cortical cells from E13.5 embryos and screened a
library of 450 kinase inhibitors for their ability to alter the proportion of *Satb2*^{tdTom} neurons at
85 day-in-vitro two (DIV2) (Grinshtein et al., 2016). APY69 was among the strongest modulators
of *Satb2* expression with its dose-dependent effect (Fig. 1b, Table S1). APY69 specifically

inhibits Ire1 α , an evolutionarily ancient bimodal transmembrane kinase and RNase, which acts as an endoplasmic reticulum (ER) stress sensor, where it catalyzes mRNA splicing in a spliceosome-independent manner (Lee et al., 2008; Tirasophon et al., 1998). Recent
90 evidence highlights an unfolded protein response (UPR)-independent developmental function of Ire1 α as regulator of cellular migration (Urrea et al., 2018). We hypothesized that stress-independent function of Ire1 α might be essential for specification of Satb2 identity in developing neurons.

We first studied the expression pattern of Ire1 α . Fluorescence in situ hybridization
95 (FISH) revealed a homogenous expression of Ire1 α throughout the developing cortex. We noted a strong developmental downregulation of Ire1 α in the cortex, with its highest levels at E12.5, both at the level of protein, and mRNA (Fig. 1c-1d). As opposed to isolated perinuclear puncta visible in the majority of cells, we observed single cells at E12.5 and at E14.5 with diffuse cytoplasmic Ire1 α mRNA FISH signal.

100 We then went on to validate the pharmacological screening results for APY69 (Fig. 1b) using a knock-out (KO) mouse model. Given ubiquitous expression of Ire1 α in many tissue types, its full KO results in overt developmental defects, associated with non-central nervous system-related phenotypes (Iwawaki et al., 2009). To carve out the role of Ire1 α in cortical development, we used the *Ire1 α ^{ff}* line, which enables Cre-dependent deletion of
105 exons 20–21 from the floxed *Ire1 α* allele. Upon expression of Cre, the kinase-extension nuclease (KEN) domain including RNase active site is deleted from the Ire1 α protein.

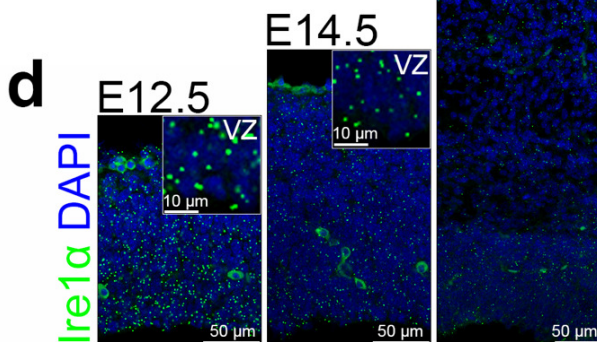
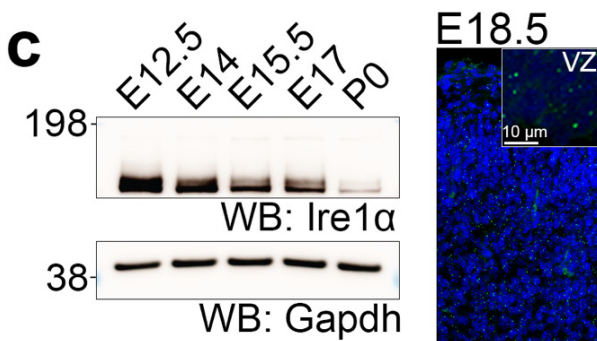
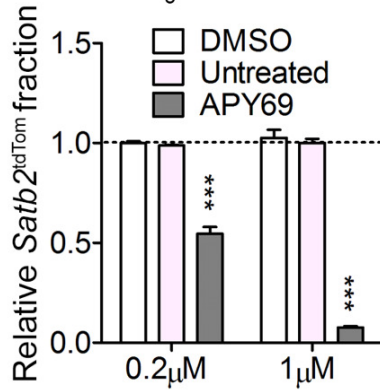
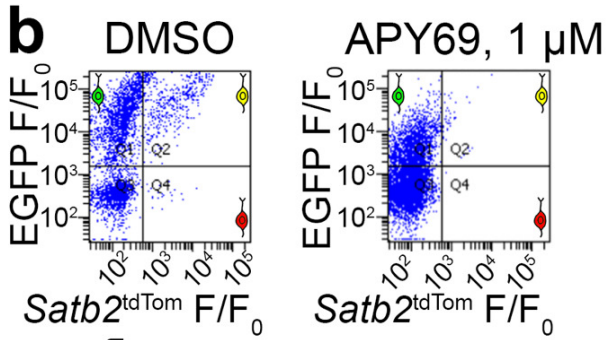
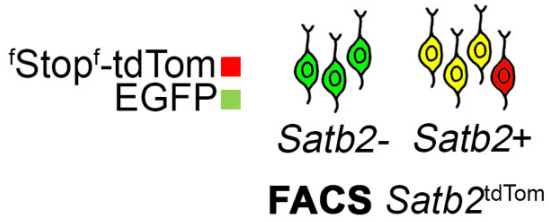
To inactivate *Ire1 α* in defined cell populations in the developing cortex, we took advantage of *in utero* electroporation (IUE), which allows for *in vivo* DNA delivery to a spatiotemporally defined subset of progenitors at the ventricular surface of the embryonic
110 cortex. Early (E12.5) neuronal progenitors display multipotency in their ability to generate both CTIP2- and Satb2-positive neuronal progeny, whereas late (E14.5) progenitors give rise to exclusively Satb2-expressing neurons populating upper layers. In our first set of experiments, we used IUE in *Ire1 α ^{ff}* mice to transfect EGFP as a control and both EGFP and Cre to induce *Ire1 α* KO in a subset of early cortical progenitors at E12.5 and derived progeny
115 (Fig. 1e). We then immunostained E16.5 coronal cortical sections for EGFP and neuronal identity markers, Satb2 and CTIP2. We first noted disrupted, highly branched morphology of *Ire1 α* KO neurons, compared to majority of bipolar control cells (Fig. 1f). Quantification revealed fewer Satb2-expressing neurons at the expense of CTIP2-positive ones in the *Ire1 α* KO (Fig. 1g), concordant with our *in vitro* results (Fig. 1b). On the contrary, overexpression
120 of human IRE1 α in wild-type early cortical progenitors at E12.5 (Fig. 1h) resulted in an increased proportion of Satb2-expressing neurons at E18.5, and remarkably, hardly any CTIP2-positive neurons (Fig. 1i).

Post-mitotic promoter NeuroD1-induced expression of Cre in *Ire1 α ^{ff}* at E12.5 did not alter the proportion of Satb2 and CTIP2 neurons at E16.5 (Fig. S1a and S1b), indicative of a
125 neuronal progenitor-embedded role of *Ire1 α* in the acquisition of neuronal identity. We observed an analogous morphological phenotype in NeuroD1-Cre expressing neurons in *Ire1 α ^{ff}* mice as in ones transfected with CAG-Cre (Fig. S1a and S1c, compare with Fig. 1f).

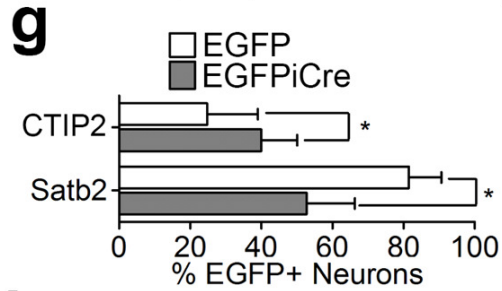
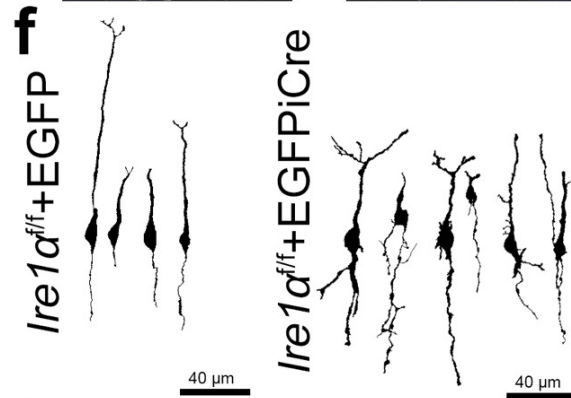
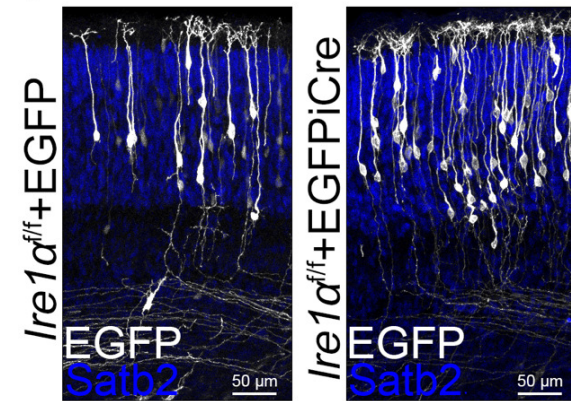
Using an *Emx1*-Cre deleter, we then conditionally inactivated *Ire1α* in the progenitors of dorsal telencephalon by establishing a *Ire1α^{fl/fl}; Emx1^{Cre/+}* mouse line, *Ire1α* cKO. We
130 injected pregnant mice with BrdU at E12.5 (Fig. S2a-S2d) and E14.5 (Fig. S2e-S2h) and quantified the proportions of *Satb2* and *CTIP2*-expressing neurons at P2, when neurogenesis is complete. Among all E12.5-derived BrdU+ cells, we counted fewer *Satb2*+ and more *CTIP2*+ cells (Fig. S2b and S2c), consistent with our IUE experiments (Fig. 1). Strikingly, in
135 E14.5 BrdU-pulsed P2 embryos, we found slightly more BrdU+/*CTIP2*+ cells (Fig. S2f and S2g), which localized to the top of the CP.

These data altogether indicate a critical requirement of *Ire1α* for the correct proportion between upper layer-associated *Satb2*-positive and deeper layer-associated *CTIP2*-expressing neurons specified during cortical development.

a Small molecule screening
in *Satb2^{Cre/+}*



e IUE E12.5 - E16.5



h IUE E12.5 - E18.5

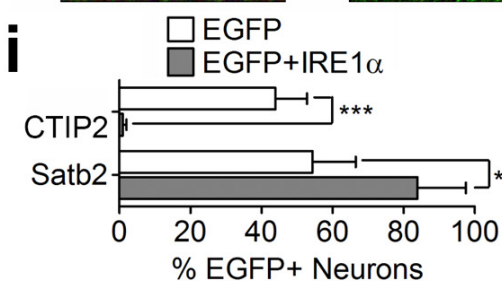
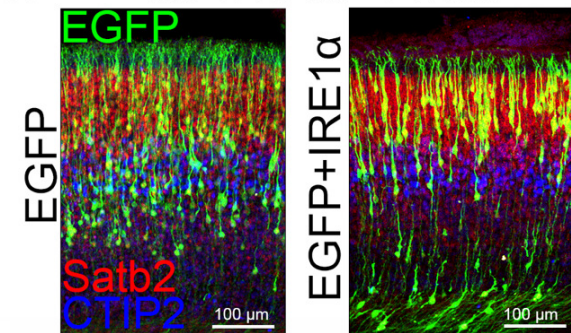


Fig. 1. Small molecule screening reveals Ire1 α activity pivotal for upper layer neuronal identity and bipolar morphology of cortical neurons.

(a) Screening workflow to identify signaling pathways upstream of *Satb2* neuronal identity. (b) Top
145 panels: representative results of flow cytometry of DIV2 neurons treated with DMSO and APY69, a
selective inhibitor of Ire1 α . Bottom panels: quantification of the proportion of *Satb2*-positive neurons.
(c) Western blotting in cortical lysates to profile developmental expression of Ire1 α . Gapdh, loading
control. (d) Fluorescence in situ hybridization for Ire1 α in cortical sections at indicated developmental
stages. Insets demonstrate enlarged fragments of the ventricular zone (VZ). (e) Representative
150 images of immunostaining against EGFP and *Satb2* in E16.5 coronal cortical sections of *Ire1 α ^{fl/fl}*
embryos after IUE at E12.5 with plasmids encoding for EGFP or EGFP and Cre simultaneously. (f)
Representative EGFP-based tracings of single neurons after IUE in (e). (g) Quantification of neuronal
cell identity after IUE described in (e). (h) Representative images of cortical coronal sections at E18.5
after IUE in wild-type E12.5 embryos to express EGFP, or EGFP and human IRE1 α . (i) Quantification
155 of neuronal cell identity after IUE described in (h). For (e), (g) and (h-i), compare Fig. S6. Bar graphs
show averages \pm S.D. Statistics for (b), one-way ANOVA with Bonferroni post-hoc test; (g) and (i),
D'Agostino-Pearson normality test and unpaired t-test. *** $p < 0.001$; $0.01 < * p < 0.05$.

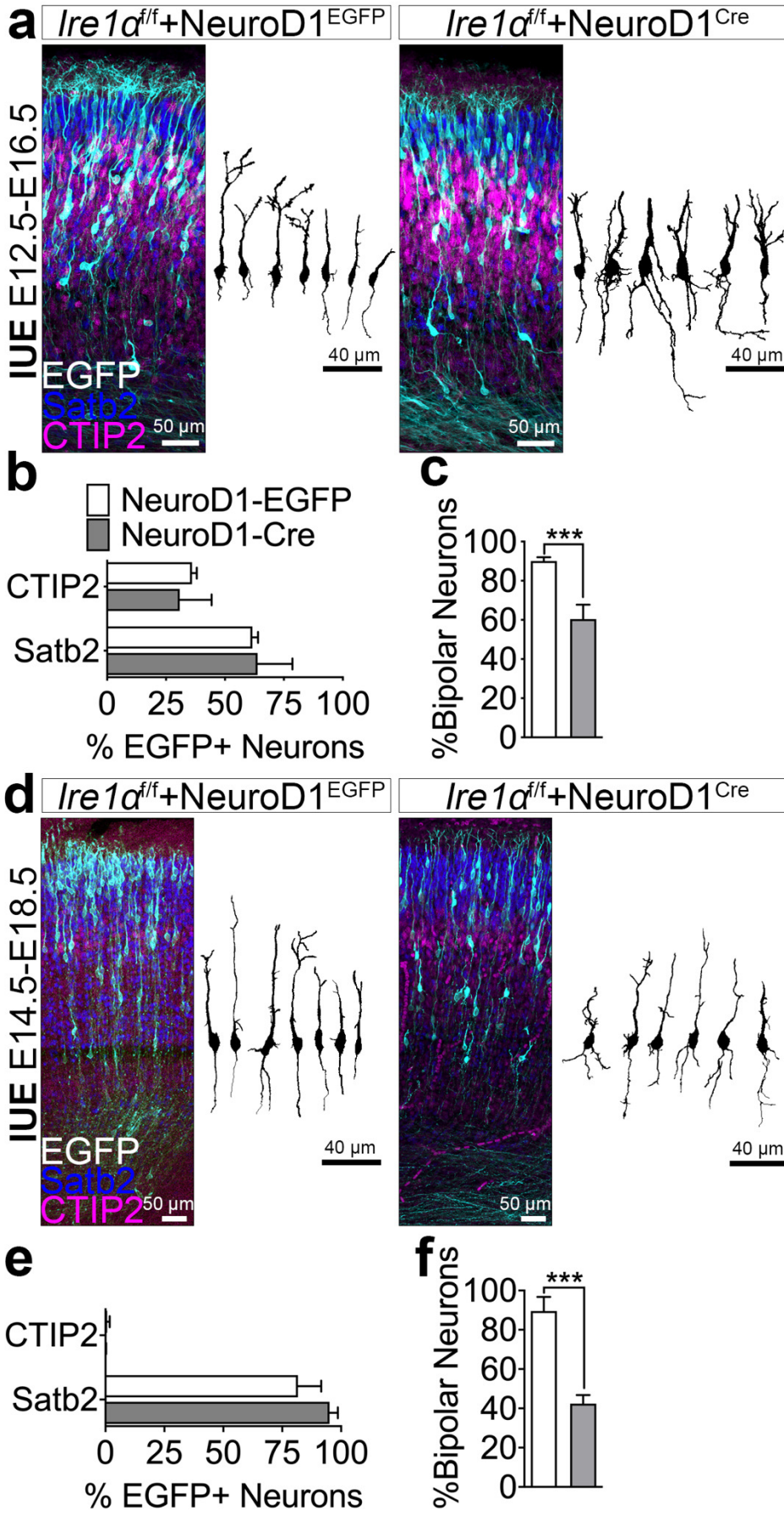
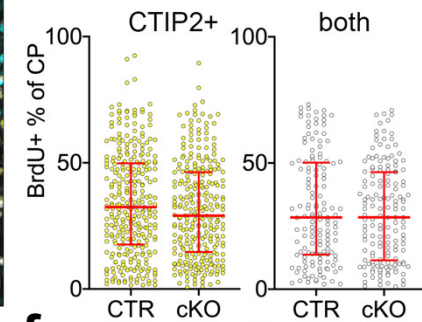
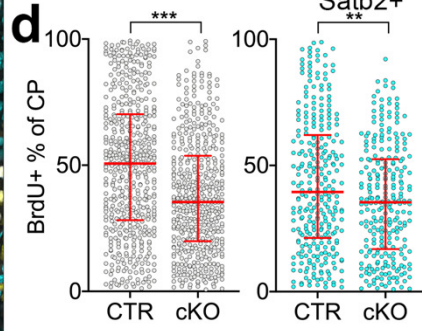
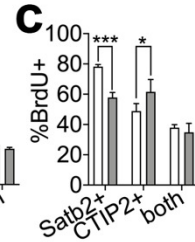
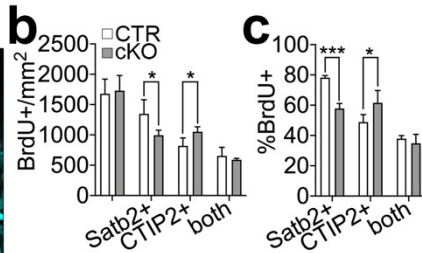
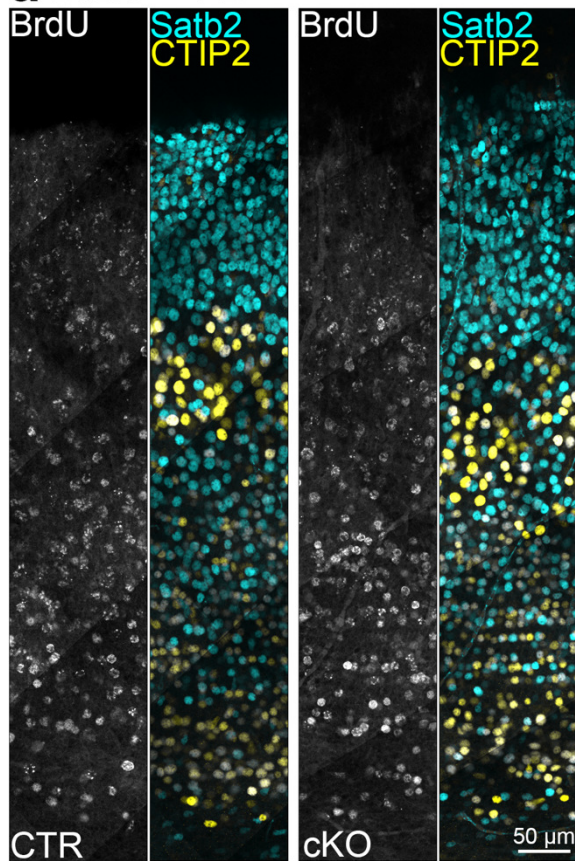


Figure S1. Related to Fig. 1. *Ire1α*-driven acquisition of the upper layer identity, but not of bipolar morphology, is embedded in neuronal progenitors.

165 (a, d) Representative images of immunostaining in coronal cortical sections from E16.5 (a) or E18.5 (d) *Ire1α^{ff}* embryos after IUE at E12.5 (a) or E14.5 (d) with plasmids encoding for EGFP under the promoter of *NeuroD1* gene or Cre under the promoter of *NeuroD1* together with loxP-Stop-loxP-EGFP. Shown are also representative, EGFP-based tracings of single neurons. (b, e) Quantification of neuronal cell identity in (a) and (d). (c, f) Average proportion of bipolar neurons in (a) and (d). Bar graphs show averages \pm S.D. For statistical analyses, D'Agostino-Pearson normality test and Mann-Whitney test. *** $p < 0.001$.

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a E12.5^{BrdU}-P2



e E14.5^{BrdU}-P2

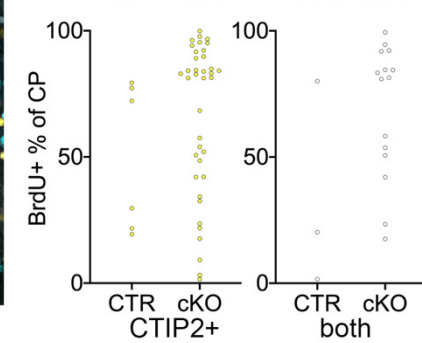
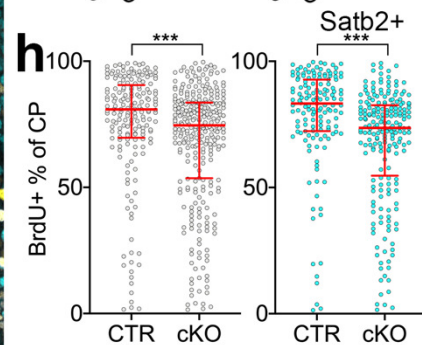
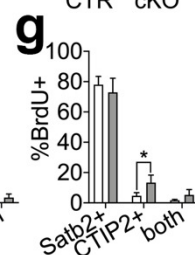
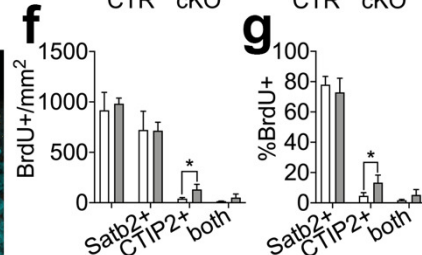
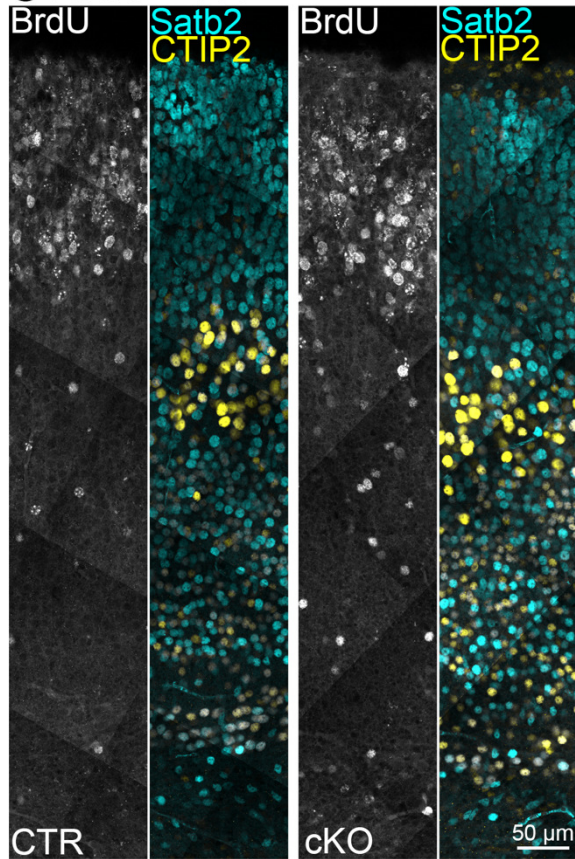


Figure S2. Related to Fig. 1. Birth dates and laminar positions of Satb2+ and CTIP2+ neurons in *Ire1α* cKO cortex.

(a, e) Representative images of immunostaining in control or cKO coronal cortical sections after BrdU injection at E12.5 (a-d) or E14.5 (e-h). (b), (f) Quantification of cell density expressing an indicated nuclear stain. "Both", cells co-expressing of Satb2 and CTIP2. (c), (g) Quantification of the proportion of neurons expressing indicated marker within all BrdU+ cells. (d), (h) Laminar distribution of neurons expressing indicated marker. The position of each neuron was normalized to the thickness of the CP and represented as a dot (100% CP – pia, 0% CP – bottom of the CP). Graph contains pooled data from indicated number of brains (Table S1). Bar graphs show averages \pm S.D. Line and error bars on (d) and (h) indicate median and interquartile range. For statistical analyses, D'Agostino-Pearson normality test and (d) and (h), Mann-Whitney test; (b-c) and (f-g) unpaired t-test. $0.001 < ** p < 0.01$; $*** p < 0.001$.

185 **Ire1 α is required for axon specification in developing cortical neurons of upper layers**

Efferent cortical connectivity is established by the correct proportion of neuronal types and specification of functional axons during development. Because of the prominent morphological phenotype for E12.5 progenitor-derived neurons (Fig. 1f), and given that
190 pyramidal upper layer neurons serve as an established model to study specification of axon-dendrite polarity (Barnes and Polleux, 2009), we electroporated E14.5 neuronal progenitors of *Ire1 α ^{fl/fl}* embryos with plasmids encoding EGFP or EGFP and Cre and fixed the brains at E18.5. In line with previously published results (Urrea et al., 2018), loss of *Ire1 α* lead to a disrupted laminar positioning of upper layer cortical neurons. While control neurons primarily
195 localized to the top of the cortical plate (CP), as predicted from their birthdate at E14.5, we noted a substantial number of Cre-expressing neurons populating the bottom of the CP as well (Fig. 2a and 2b). Consistently, in the cKO brains, both E12.5 and E14.5 BrdU-pulsed *Satb2*-expressing cells localized to the lower portions of the CP, as compared to the control cells (Fig. S2). Using expansion microscopy (ExM), we detected multiple trailing processes
200 (TPs) and many short processes originating from the soma in *Ire1 α* KO upper layer neurons (Fig. 2c). This contrasted with the majority of control neurons characterized with a single TP and lack of additional neurites emanating from the soma at this stage (Fig. 2d). Disruption of bipolar morphology and aberrant laminar positioning within the CP was independent of *Satb2* expression (Fig. 2e). We also observed morphological defects induced by IUE of *NeuroD1*-
205 Cre inducing *Ire1 α* KO in differentiating neurons (Fig. S1d-S1f). These data indicate *Ire1 α* has both a progenitor and post-mitotic neuron embedded role in the regulation of neuronal morphology during cortex development.

To study the identity of excessive processes specified by *Ire1 α* KO neurons, we took advantage of *ex utero* electroporation (EUE), which permits modulation of gene expression
210 in the neuronal progenitors followed by *in vitro* primary neuronal culture. We expressed EGFP or simultaneously EGFP and Cre in E14.5 upper layer neuron progenitors of *Ire1 α ^{fl/fl}* embryonic cortices and cultured isolated primary cells until DIV4, when axons are established in control neurons (Ambrozkiwicz et al., 2018). As expected, the majority of control cells projected a single longest neurite, while *Ire1 α* KO cells were symmetrical and failed to project
215 a long axon (Fig. 2f). We then immunostained the fixed cell cultures for axonal and dendritic markers (*Tau-1* and *MAP2*, respectively) and counted the number of axons projected by single neurons (Fig. 2g). We failed to identify an axon in a fraction of *Ire1 α* KO neurons, and identified a significant proportion of neurons that projected multiple axons, as compared to polarized control cells (Fig. 2h). Where present, *Ire1 α* KO axons were shorter (Fig. 2i) and
220 more branched (Fig. 2j), as compared to control ones. *Ire1 α* KO neurons with no axons exhibited increased dendritic branching, possibly implicating an axon-to-dendrite identity switch of single neurites. Axon possessing *Ire1 α* KO neurons exhibited a normal dendritic tree, when compared to controls (Fig. 2k and 2l). This indicates formation of excess axons upon *Ire1 α* KO, rather than a dendrite-to-axon transition in neurons with multiple axons.
225 Morphological defects in the KO were associated with somatic distribution of ER tubules visualized by EGFP-*Sec61 β* expression as compared to its polarized localization in neurites

of control cells (Fig. 2m and 2n). This indicates a possible involvement of $Ire1\alpha$ in ER-driven neuronal polarization.

We then asked if such profound morphological defects have consequences on axon-
230 related neurophysiology. At DIV6, axons of upper layer control neurons are enriched for Tau-
1 (Fig. S3a), Ankyrin G (Fig. S3b), and voltage-gated Na^+ channels (Na_v s; Fig. S3c). In *Ire1\alpha*
KO neurons, we observed a perinuclear somatic localization of these axonal markers,
indicative of their aberrant subcellular localization. Current clamp recordings from *Ire1\alpha^{fl/fl}*
autaptic hippocampal neurons infected with lentiviruses encoding for EGFP or Cre showed a
235 higher number of action potentials (APs) generated upon lower current injections (Fig. S3d
and S3e). This might be attributable to more axon initial segments (AIS) specified by *Ire1\alpha*
KO neurons and/or their altered molecular composition, which increases the chance of firing
at lower currents. Importantly, overall membrane integrity and conductance of *Ire1\alpha* KO
neurons were unaltered (Fig. S3f and S3g). Altogether this evidence reveals that loss of
240 bipolar morphology in developing *Ire1\alpha* KO neurons is associated with defects in axon
specification, subcellular mislocalization of axonal markers and increased current sensitivity.

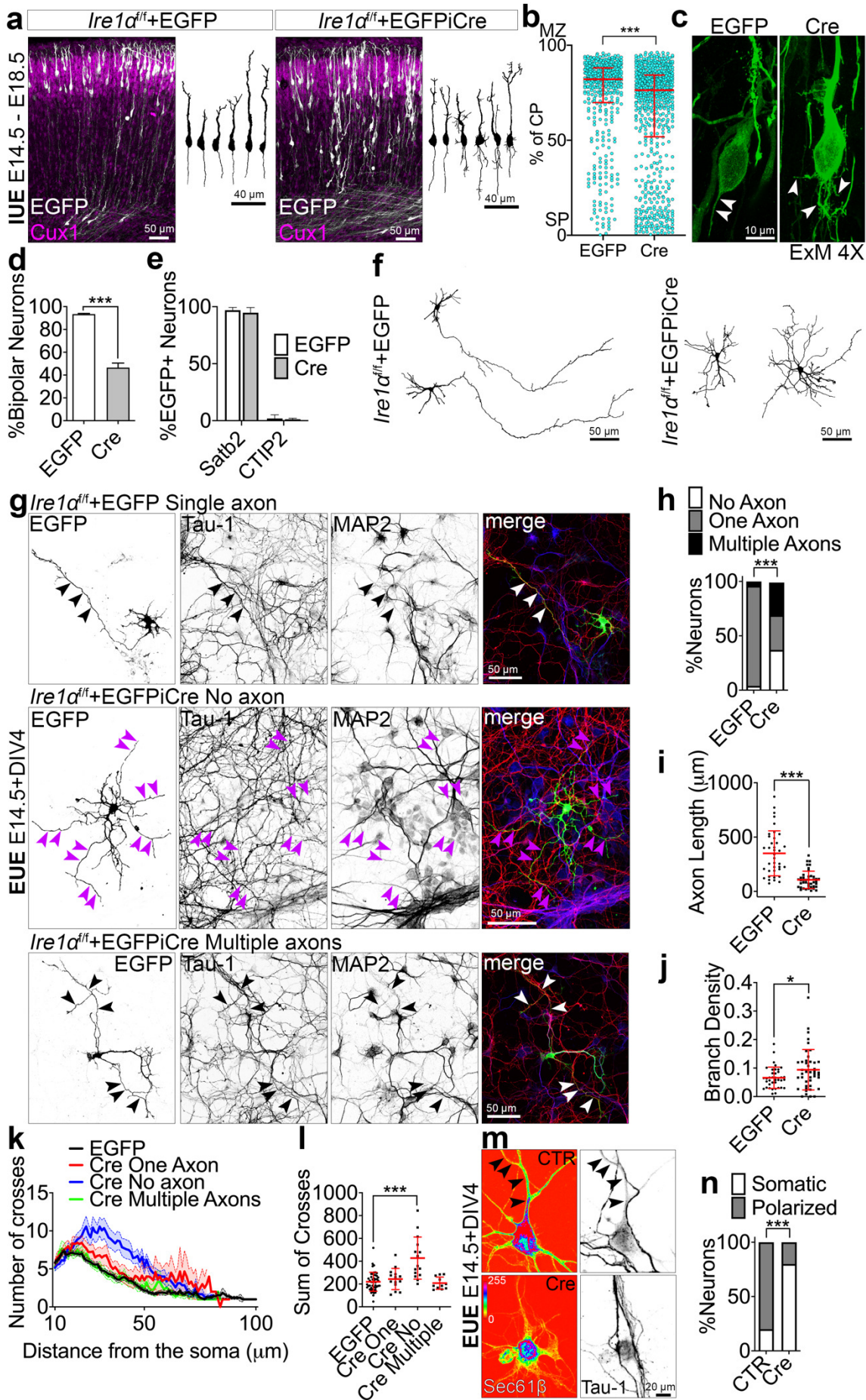


Fig. 2. *Ire1α* is indispensable for bipolar morphology, axon specification and proper laminar positioning of upper layer neurons.

(a) Representative images of EGFP and upper layer-expressed Cux1 immunostaining in coronal cortical sections from e18.5 *Ire1α^{ff}* embryos after IUE at e14.5 with indicated plasmids and semi-automatic, EGFP-fluorescence based tracings of single neurons. (b) Laminar distribution of cortical neurons in e18.5 *Ire1α^{ff}* brains after IUE described in (a). The position of each neuron was normalized to the thickness of the CP and represented as a dot. 0% - Subplate (SP), 100% - Marginal Zone (MZ). Graph contains pooled data from indicated number of brains (Table S1). (c) Representative images of EGFP immunostaining after expansion microscopy (ExM) in cortical sections from (a). Arrowheads point to trailing processes. (d) Average proportion of bipolar neurons from (a). (e) Quantification of neuronal identities in experiment in (a). (f) Representative EGFP-based tracings of *Ire1α^{ff}* neurons at DIV4 after EUE to express EGFP or EGFP and Cre. (g) Neurons in (f) were fixed and immunostained with axonal or dendritic markers. White and black arrowheads indicate axons, purple ones indicate neurites lacking Tau-1 expression. (h) The number of axons projected from a single neuron. Average axon length (i), average axon branch density per micrometer (j), Sholl analysis diagram (k) and average sum of crossing dendrites (l) in neurons expressing EGFP or EGFP and Cre, projecting one, or multiple axons. (m) Representative EGFP signals and Tau-1 immunostaining of *Ire1α^{ff}* DIV4 neurons after EUE to co-express EGFP-Sec61β and empty vector (CTR) or Cre. (n) Quantification of EGFP-Sec61β localization in experiment in (m). Line and error bars on (b) indicate median and interquartile range, scatter plots in (i), (j), (l) and bar graphs on (d) and (e) show averages ± S.D. Results on (k) are represented as averages ± S.E.M. Statistics for (b), (d-e), (i-j), D'Agostino-Pearson normality test and Mann-Whitney or unpaired t-test; (h), Chi-square test; (l), Kruskal-Wallis with Dunn's multiple comparisons test; (n), Fisher's test. *** p < 0.001; 0.01 < * p < 0.05.

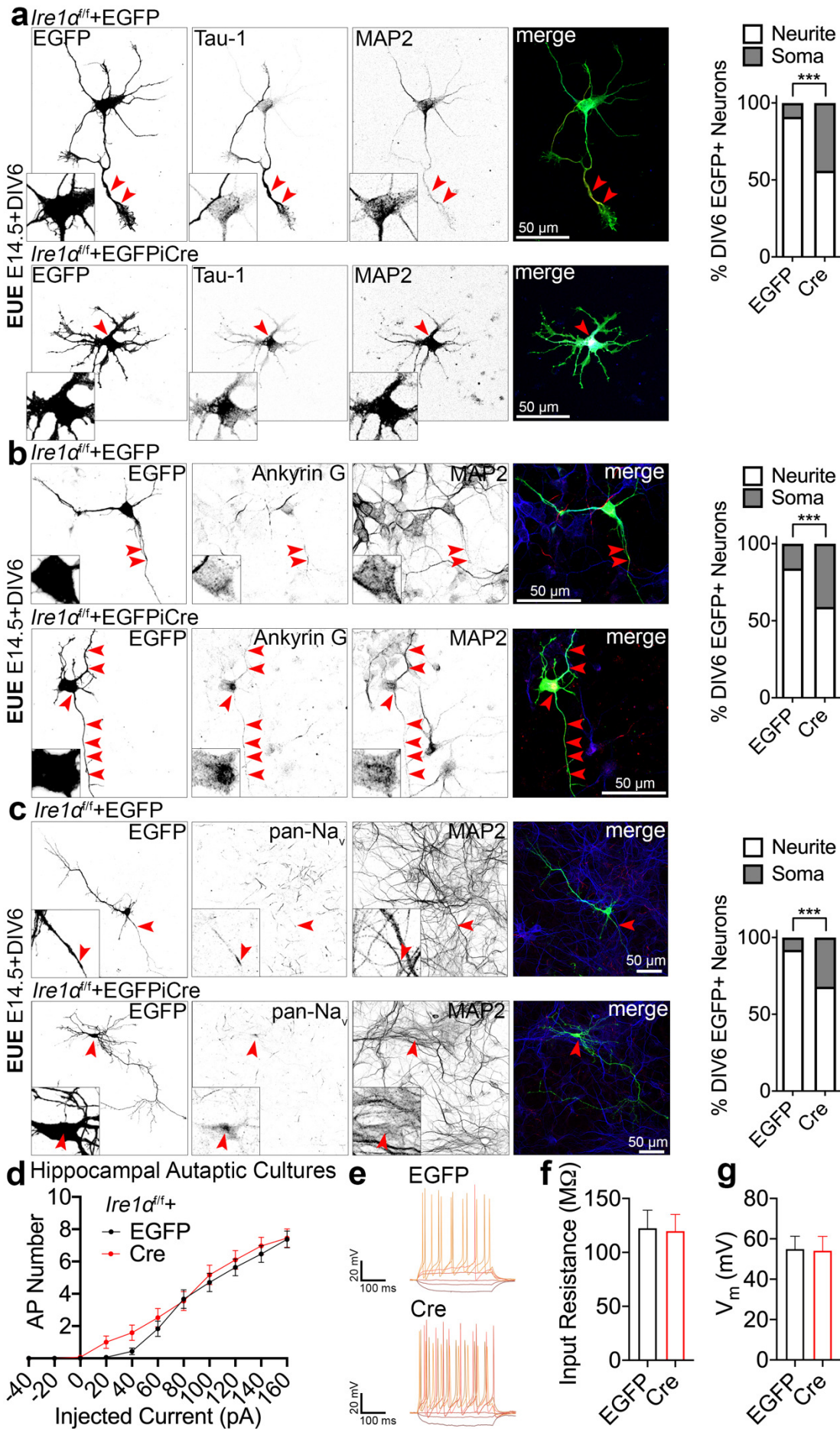


Fig. S3. Related to Fig. 2. Loss of *Ire1α* disrupts axon initial segment and current responses.

270 (a-c) Images of representative DIV6 primary cortical *Ire1α^{ff}* neurons after EUE at E14.5 for EGFP or
EGFP and Cre expression. Neuronal cultures were immunolabeled for indicated axonal and dendritic
marker proteins. Graphs show quantification of the subcellular distribution of axonal markers. *Ire1α*-
deficient neurons display somatic accumulations of Tau-1, Ankyrin G and Nav. Red arrowheads
indicate enrichment of axonal markers. Insets on the bottom left of black and white images are zoom-
275 ins to the somata or places of axonal marker accumulation. (d-g) Electrophysiological recordings from
autaptic hippocampal cultures prepared from P0 *Ire1α^{ff}* pups. Neuronal cultures were infected with
lentiviruses encoding for EGFP or Cre. (d) Quantification of the number of APs generated after
injection of increasing amount of current. (e) Example traces of recordings described in (d). Average
input resistance (f) and holding potential (g). Data points and error bars on (d) represent mean ±
280 S.E.M. and graphs on (f-g), mean ± S.D. For statistical analyses, (a-c), Fisher's test; (d), one-way
ANOVA; (f-g), Mann-Whitney test. *** p < 0.001.

Ire1 α drives upper layer specification and axon formation in the neocortex by controlling expression levels of translation regulators

285 During the UPR, robust cellular reprogramming is driven by Ire1 α signaling network (Cox et al., 1993; Mori et al., 1993). Ire1 α activation is paralleled by suppression of general translation (Di Prisco et al., 2014). We hypothesized that also during development, Ire1 α orchestrates upper layer subtype and axonal polarization by influencing protein translation and thereby affecting expression of key developmental determinants. To test this, we analyzed the
290 ribosome profile in forebrain-specific *Ire1 α* cKO, and we purified actively translating ribosomes *ex vivo* (Kraushar et al., 2021). Given a scarcity of material at early embryonic stages, we used E18.5 brains. Sucrose density gradients revealed higher level of polysomes in *Ire1 α* cKO cortices (Fig. 3a).

We then investigated which RNAs localize to polysome fraction in *Ire1 α* cKO cortex.
295 We performed RNA sequencing in bulk cortex of control and cKO as well as in the polysome fraction purified from the control and *Ire1 α* cKO cortices. Notably, we did not detect gross changes in the level of the transcriptome in the bulk RNAseq [(Fig. 3b; cutoff at corrected p-value 0.05, AY036118 lncRNA outlier comes from inflated expression counts given its alignment with *Rn45s* repeat (Kimmel et al., 2019)]. Polysome-associated RNA in *Ire1 α* cKO
300 cortices (depicted as colored data points, cutoff at corrected p-value 0.05) encoded for structural components of the ribosome, proteins acting on RNA, revealed by the gene set enrichment analysis (GSEA, Fig. 3d), and other molecules regulating translation and initiation factors (Table S2). These results indicate that *Ire1 α* loss in the cortex alters translation of RNAs crucial for ribosome complex function.

305 We then investigated the expression levels of canonical translation regulators using Western blotting and E18.5 cortical lysates (Fig. S4a and S4b). We detected no gross changes in the level of classical regulators of translation in the developing cortex, like S6-P, Fragile X Mental Retardation Protein 1 (FMRP), or eukaryotic initiation factor 2 α . Notably, the level of spliced Xbp1, a cellular stress-associated Ire1 α substrate was not altered, indicating
310 a non-canonical UPR-independent role of Ire1 α in the regulation of polysome level. We identified upregulated levels of eukaryotic elongation factor 2 (eEF-2), downregulated phosphorylated eEF2 on Thr56 and decreased level of eukaryotic initiation factor 4A1 (eIF4A1) in the *Ire1 α* cKO cortex (Fig. 3e and 3f). Given no gross changes on the RNA level (Fig. 3b), we conclude that the expression level changes for these factors are of translational
315 nature.

To mimic the cKO scenario, we then overexpressed eEF-2 using IUE in e12.5 and e14.5 wild type embryos (Fig. S4c-S4g). Four days later, we detected no alteration of Satb2 proportion in the neuronal progeny after eEF-2 OE (Fig. S4d and S4f). However, we did detect a positioning defect of upper layer neurons, similar to the one observed upon loss of *Ire1 α*
320 (Fig. S4g). Next, we inactivated *eIF4A1* using CRISPR-Cas9 technology (Fig. 3g), which allows for simultaneous sgRNA and humanized Cas9 nickase delivery to developing cortex using IUE in e12.5 wild type embryos (Shinmyo et al., 2016). The loss of *eIF4A1* resembled *Ire1 α* KO (Fig. 1e and 1g) regarding the types of neurons generated, with less Satb2-expressing neurons, at the expense of CTIP2-positive cells (Fig. 3h). Loss of *eIF4A1* from
325 upper layer progenitors did not affect laminar positioning in the cortex at e18.5 (Fig. S4e-S4g). Both eIF4A1 and eEF-2 were highly and homogeneously expressed in the E12.5 cortex,

and downregulated at E14.5, resembling the expression pattern of *Ire1α* (Fig. S4h-S4i, compare Fig. 1d). We also studied the polarity acquisition in DIV4 neurons after EUE-induced eEF-2 OE, *eIF4A1* KO and simultaneous eEF-2 OE and *eIF4A1* KO (Fig. S4j-S4k). We found
330 that all three manipulations lead to emergence of neurons with 0 axons, characterized by absence of Tau-1 enrichment in a single neurite, partially reminiscent of *Ire1α* KO neurons (Fig. 2g-2h). Notably, simultaneous eEF-2 OE and *eIF4A1* KO resulted in the highest proportion of neurons without an axon.

Altogether, these findings demonstrate that *Ire1α* controls expression levels of
335 translation regulators. In particular, initiation factor eIF4A1 acts downstream of *Ire1α* to control *Satb2* expression and thereby neuronal subtype identity determination in the developing neocortex, while elongation factor eEF-2 regulates the laminar distribution of upper layer neurons. Tightly regulated levels of both eIF4A1 and eEF-2 in neuronal progenitors are critical for specification of an axon in developing neurons.

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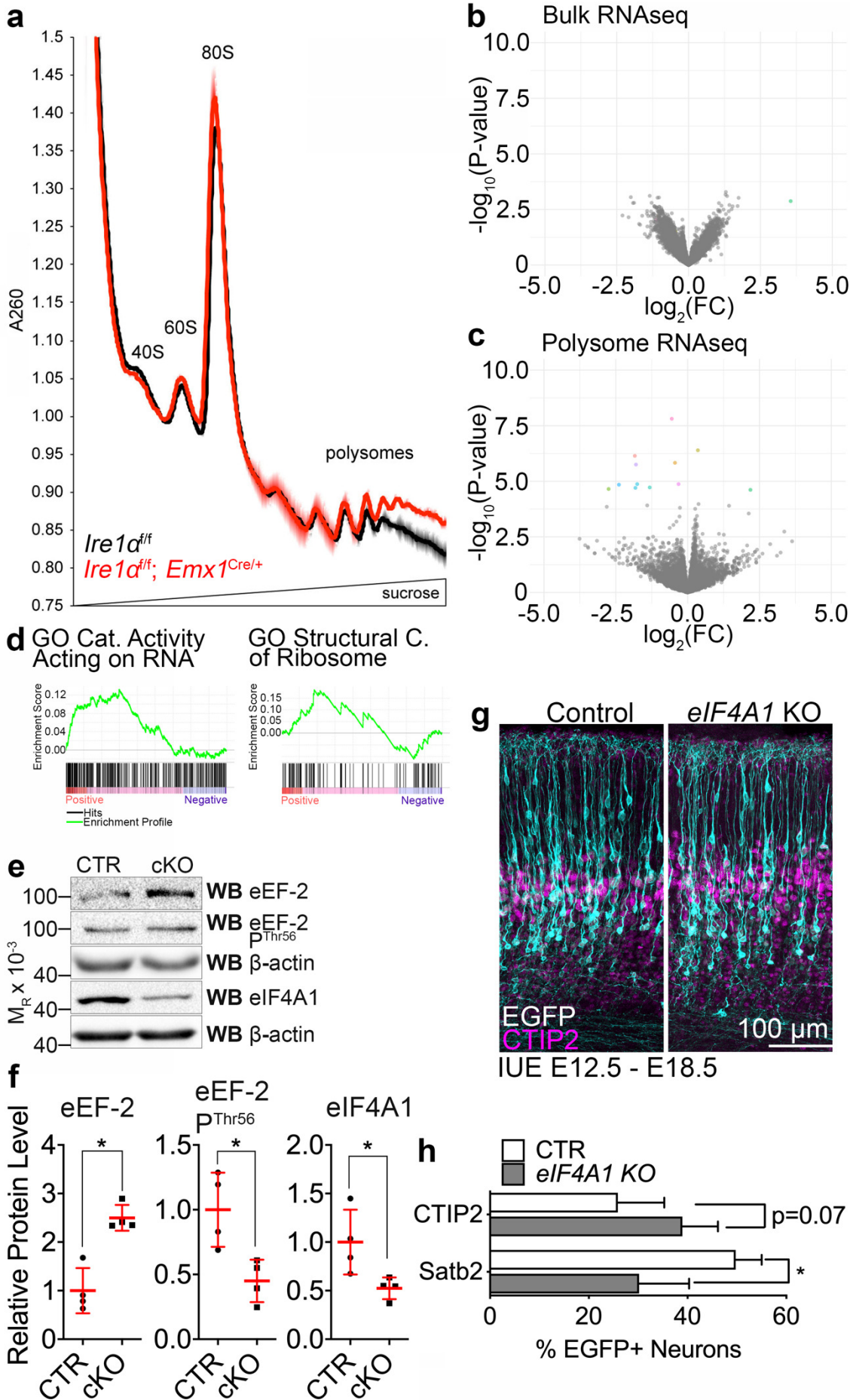
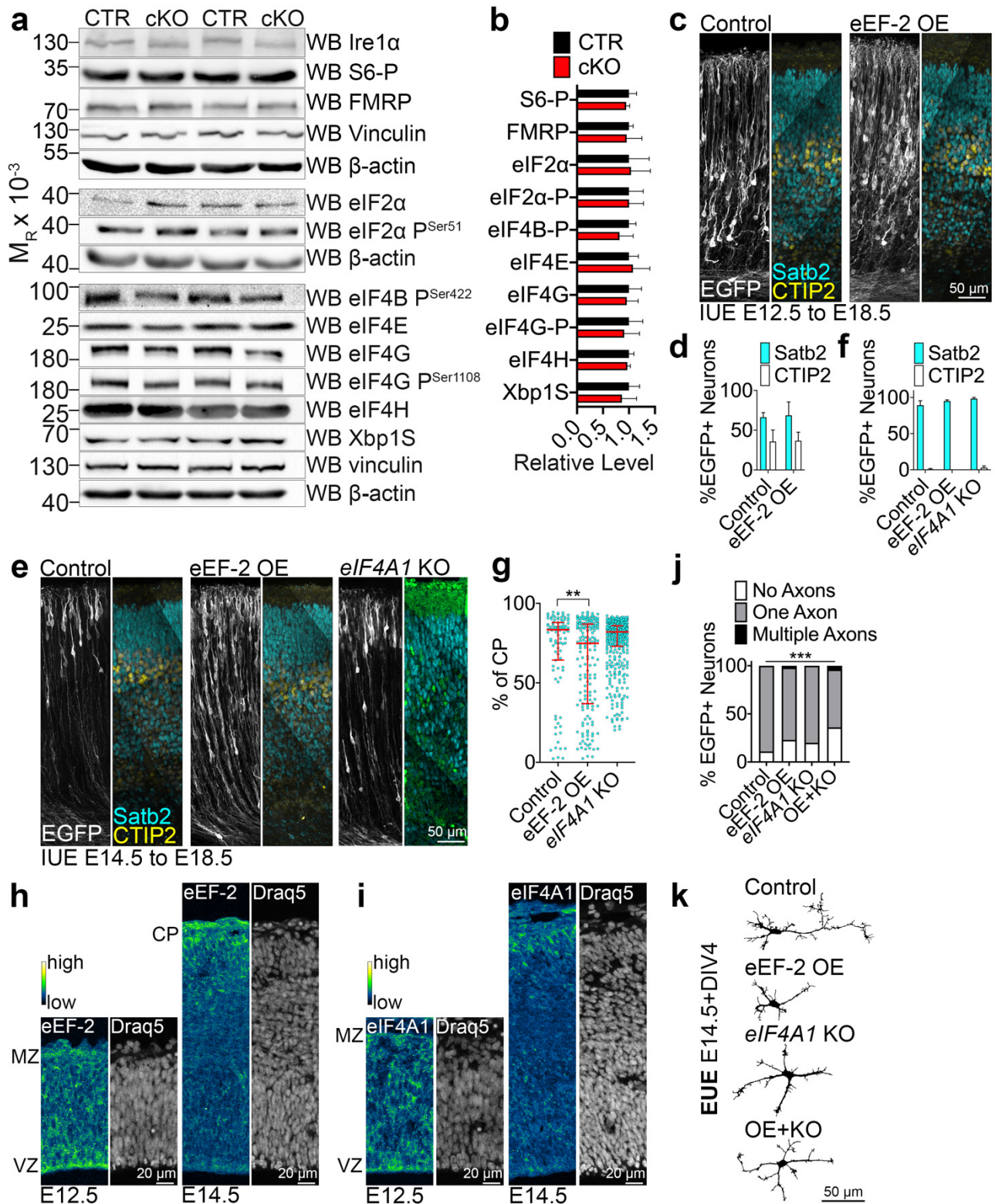


Fig. 3. Ire1 α -mediated protein translation regulation drives upper layer identity acquisition in the cortex.

(a) Analytic density gradient fractionation of A260-normalized E18.5 neocortex lysates, measuring the relative abundance of ribosomal subunits, 80S ribosomes, and polysomes. A260 curves plotted as mean \pm S.D. across replicate fractionations (n = 2, each comprising 7-8 brains per condition), baseline (1.0) centered at onset of 40S peak. (b and c) Volcano plots for *Ire1 α* cKO RNAseq in bulk tissue (b) and polysome-enriched RNA (c). Colored data points represent significantly regulated genes. FC; fold change. All P-values are Benjamini-Hochburg adjusted p-values. (d) Top two GSEA enrichment plots for cKO polysome RNA fraction versus control polysome RNA fraction, Catalytic Activity Acting on RNA and Structural Component of Ribosome (Table S2). (e) Representative Western blotting results using the control and *Ire1 α* cKO E18.5 cortical lysates. (f) Quantifications of the protein level from (e). (g) Representative images of immunostaining against EGFP and CTIP2 in E18.5 coronal cortical sections of wild type embryos after IUE at E12.5 with gRNAs and Cas9 nickase to achieve indicated genotypes. (h) Quantification of neuronal cell identity in the experiment in (g). Compare Fig. S6. Red line and error bars on (f) and bar graphs of (h) indicate mean \pm S.D. Statistics for (f) and (h), D'Agostino-Pearson normality test and Mann-Whitney test. 0.01 < * p < 0.05.



360

Fig. S4. Related to Fig. 3. *Ire1α*-mediated protein translation regulation employs stress-independent signaling pathways and drives axon specification.

(a) Representative results of Western blotting in E18.5 cortical lysates from control and *Ire1α* cKO using indicated antibodies. (b) The expression level of analyzed protein was normalized to the amount of beta-actin in the sample and expressed as relative protein level. (c-g) Representative images (stitched tiles) of EGFP fluorescence signals and immunolabeling for Satb2 and CTIP in E18.5 brains

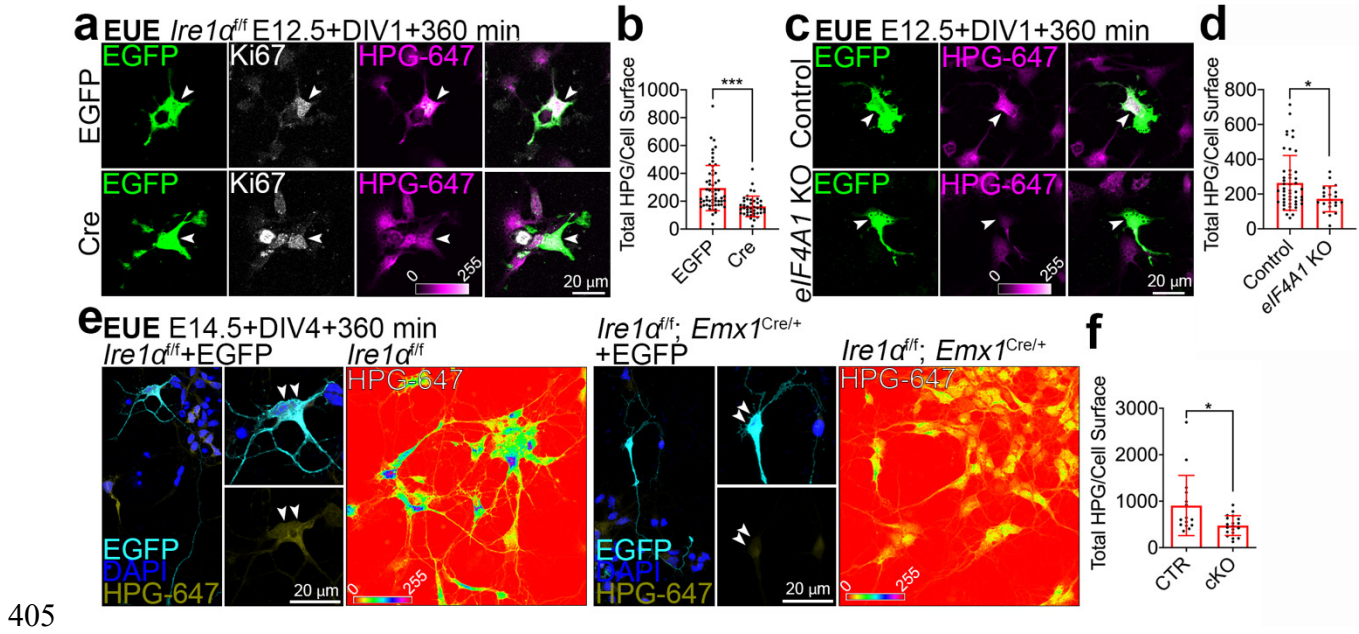
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of wild type embryos after IUE at E12.5 (c-d), or E14.5 (e-g) with plasmids encoding for EGFP (Control) or for EGFP and eEF-2 (eEF-2 OE), or for EGFP and eIF4A1 gRNAs and Cas9 nickase (*eIF4A1* KO). Quantification of neuronal identity (d and f) and positioning within the cortical plate (g).
370 (h-i) Representative images of the immunostaining in the coronal cortical slice for indicated proteins and Draq5, a nuclear marker. VZ, ventricular zone; MZ, marginal zone; CP, cortical plate. Graph contains pooled data from indicated number of brains (Table S1). (j) The number of axons projected from a single neuron at DIV4 after EUE to achieve indicated genotypes. Neurons were fixed and immunolabeled from axonal and dendritic markers (compare with Fig. 2). OE+KO, simultaneous eEF-
375 2 OE and eIF4A1 KO. (k) Representative EGFP-based tracings of DIV4 neurons from (j). Apart from Control, shown are neurons with 0 axons. Bar graphs indicate mean \pm S.D. Line and error bars on (g) indicate median and interquartile range and scatter plots individual positions of single neurons. For statistics on (b) unpaired t-test with post-hoc Holm-Sidak for multiple comparisons; (d), Mann-Whitney test; (f), (g), Kruskal-Wallis test with Dunn's multiple comparisons; (j), Chi-square test. $0.001 < ** p <$
380 0.01 ; $*** p < 0.001$.

Ire1 α is a regulator of protein translation rates in the developing cortex

Increased polysome level in the *Ire1 α* cKO may represent ribosomes elongating more slowly or stalling, and thus accumulating in the heavy fraction (Eshraghi et al., 2021). To test this
385 hypothesis, we examined translation rates in *Ire1 α* cKO neurons using FUNCAT (Beatty and
Tirrell, 2008; Dieterich et al., 2010). The method relies on replacing L-methionine with its
alkyne analog L-homopropargylglycine (HPG) in the medium of primary cultured cells, and
HPG detection in a Huisgen alkyne-azide cycloaddition reaction. Within a cell surface area,
the fluorescence intensity of azide-coupled Alexa-647 is proportional to the amount of
390 incorporated HPG into newly synthesized proteins in defined time and serves as an estimate
of translation rate in a given cell (Fig. 4).

We focused our analyses on a uniform cell population to circumvent a possibility of
confounding effects of varying translation rates in different cell types. Control of neuronal
identity mediated by *Ire1 α* is embedded in early neuronal progenitors (Fig. 1g and S1b). For
395 this reason, we first investigated protein synthesis rates using EUE to induce *Ire1 α* loss in
early E12.5 multipotent neuronal progenitors. At DIV1, we quantified lower translation rates
in Cre-expressing mitotic marker Ki67-positive (Scholzen et al., 2002) *Ire1 α* KO progenitors
as compared to EGFP-expressing *Ire1 α ^{fl/fl}* ones (Fig. 4a). Similarly, we detected lower
translation rates in E12.5 mitotic progenitors upon loss of *eIF4A1* (Fig. 4c and 4d). We
400 quantified approximately 50% lower rates of translation in DIV4 upper layer *Ire1 α* cKO
neurons born from E14.5 cortical progenitors as compared to control cells (Fig. 4e and 4f).
Taken together, regardless of the cell type, *Ire1 α* loss engenders a decrease in the rate of
protein synthesis. In the light of these data, *Ire1 α* loss-mediated increase in polysome level
may represent stalled ribosomes, accumulated in the polysome fraction.



405

Fig. 4. Loss of *Ire1α* results in lower translation rates.

(a), (c), (e) Images of representative primary cortical neurons prepared from *Ire1α^{fl/fl}* (a), wild type (c) or control and *Ire1α* cKO (e) embryos after EUE at E12.5 (a and c) or E14.5 (e) with indicated plasmids. Neurons were fed L-homopargylglycine (HPG) for 360 min prior to fixation at DIV1 or 410 DIV4. HPG was detected with Sulfo-Cyanine5 azide. (e) Right panels: images of HPG incorporation in control and cKO primary DIV4 neurons prepared from E14.5 cortex. White arrowheads point to Ki67-positive progenitors (a and c) or to somata of neurons derived from E14.5 progenitors (e). (b), (d), (f) Quantification of HPG incorporation. Bar graph represent data points for every analyzed cell and averages \pm S.D. Statistics for (b), (d), and (f), D'Agostino-Pearson normality test and Mann-Whitney test; $0.01 < * p < 0.05$; $*** p < 0.001$.

Deeper layer cortical neurons exhibit higher translation rates than upper layer neurons

Given lower translation rates in *Ire1α* cKO neurons (Fig. 4), and our findings of *Ire1α*-mediated specification of neuronal subtype (Fig. 1), we next asked whether differences in translation rates are an intrinsic feature of neurons of different cortical layers.

In order to test this, on the same day we nucleofected two populations of e12 and e14 primary cortical cells with dsRed and EGFP-expressing plasmids, respectively, mixed the now differentially labeled cells, and cultured together to tightly control for the microenvironment (to separate these sets of experiments from ones performed on *Ire1α*, we labeled the embryonic stages with a lowercase “e”). We then fixed the cells at DIV1 and DIV5, following an HPG pulse of varying duration (Fig. 5). First, we validated what cell types are present at these two culture stages using immunostaining against proliferative markers Ki67 and Pax6, and postmitotic identity markers (Fig. 5a and 5b). Our nucleofection and culture system enriches for deeper layer marker-expressing neurons derived from e12 cortex, for Satb2 in neurons prepared from e14 cortex, and generally for postmitotic neurons (Fig. 5b), with only a minor fraction of cycling cells or radial progenitors still present at DIV1 (Fig. 5c). We pulsed HPG in the methionine-free cell culture medium for 60, 120 and 240 minutes at DIV1 (for immature neurons) and at DIV5 (Fig. 4d). At DIV1, following HPG application, we quantified significantly higher HPG incorporation level in e12- (dsRed+) versus e14- (EGFP+) derived neurons (Fig. 5e and 5f). We also detected higher HPG incorporation rates at DIV5, with a milder difference between the two cell populations. Secondly, we observed, that in both e12- and e14-derived neuronal cultures, HPG incorporation rates were higher at DIV1 when compared to DIV5 (Fig. 5e and 5g). The addition of cycloheximide (CHX), a translation inhibitor (Schneider-Poetsch et al., 2010), attenuates HPG incorporation (Fig. 5h), reinforcing the translation-specific nature of our findings. Altogether, these data indicate that deep layer-fated neurons have inherently higher translation rates compared to upper layer neurons.

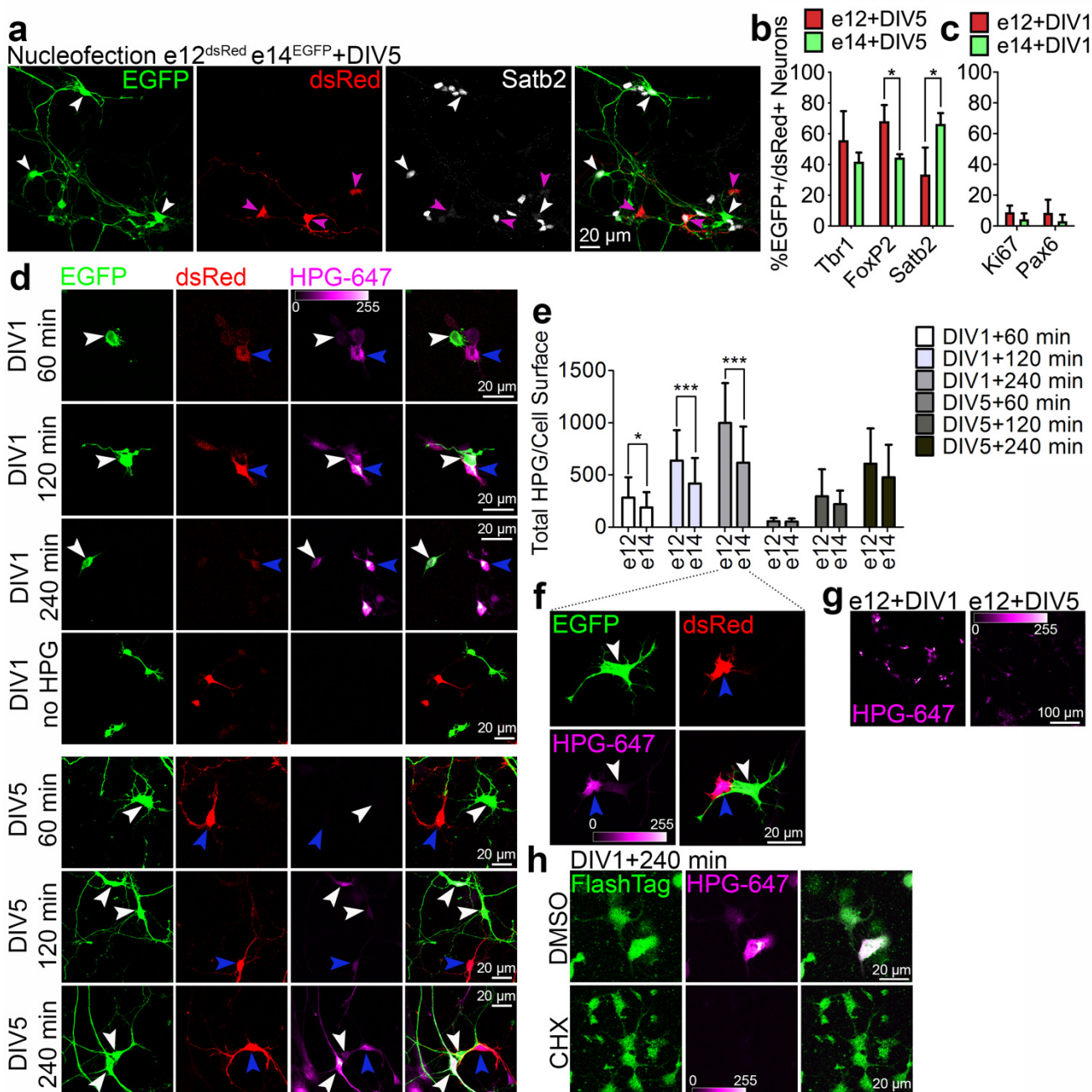


Fig. 5. Increased translation rate is a hallmark of early born, deeper layer postmitotic neurons
 445 **in the developing cortex.** (a) Images of primary DIV5 neurons immunolabeled for EGFP, dsRed,
 and Satb2. Cortical cells from e12 embryo were nucleofected to express dsRed (purple arrowheads),
 and cortical cells from e14 embryo to express EGFP (white arrowheads). Both cell populations were
 mixed together and plated on a single glass coverslip. (b, c) Quantification of the cell identity in DIV5
 neurons (b) and DIV1 primary cells (c) derived from e12 or e14 cortex. (d) Images of primary cortical
 450 cells immunolabeled for EGFP (white arrowheads) and dsRed (blue arrowheads), pulsed with HPG
 for indicated amount of time before fixation at DIV1 and DIV5. HPG was detected with far red
 fluorophore-coupled azide. (e) Incorporation of HPG was quantified as intensity of fluorescence
 signal normalized to the cell surface in primary neurons. (f) DIV1 primary cortical cells from (a),
 fed with HPG for 240 min. Note significantly higher HPG uptake in e12 cortex-derived immature
 neurons. (g) HPG labeling in primary neurons at DIV1 and DIV5 prepared from e12 cortex. Note
 455 significantly higher HPG uptake in primary cultures at DIV1. (h) DIV1 primary cortical cells
 from e14 cortex, fed with HPG

for 240 min in presence of DMSO or 10 $\mu\text{g}/\text{mL}$ cycloheximide (CHX). Note strongly reduced HPG labeling in cells cultured with CHX. FlashTag was added to the medium to label all cells. Bar graphs indicate mean \pm S.D. Statistics for (b) and (c), D'Agostino-Pearson normality test and Mann-Whitney 460 or unpaired t-test; (e), two-way ANOVA with Bonferroni post-tests. *** $p < 0.001$; $0.01 < * p < 0.05$.

The transition from multipotent neuronal progenitors to progenitors of upper layer neurons is associated with a pronounced upregulation of translation

Next, we studied the global translation rates in cycling progenitors of deeper and upper layer
465 neurons and in their directly derived progeny. To address this, we used EUE, which unlike
the nucleofection (Fig. 5) allows for labeling cycling progenitors. We transfected ventricular
e12 progenitors with dsRed- and e14 progenitors with EGFP-encoding vectors, then cultured
the cells until DIV1 and DIV5, with HPG-feeding prior to fixation. We then quantified HPG
incorporation in Ki67-labeled DIV1 progenitors and in DIV5 postmitotic neurons (Fig. 6a) and
470 found that e14 upper layer progenitors display profoundly higher translation rates as
compared to e12 multipotent progenitors. Additionally, consistently with results shown in Fig.
5, e12 progenitor-derived postmitotic neurons display higher HPG incorporation rates as
compared to e14 progenitor-derived cells. Translation rates undergo a significant
upregulation in e14 cortical progenitors that dramatically decreases during postmitotic
475 differentiation of an upper layer neuron (Fig. 6b). The majority of e12 progenitor-derived
postmitotic cells co-express both Satb2 and CTIP2 in culture, and e14 progenitor-derived
ones express Satb2, but not CTIP2 (Fig. 6c and 6d).

Taken together, the translation rate is a dynamic feature of cortical progenitors and
their derived progeny, and likely represents cell- and stage-specific requirements of different
480 protein sets during development.

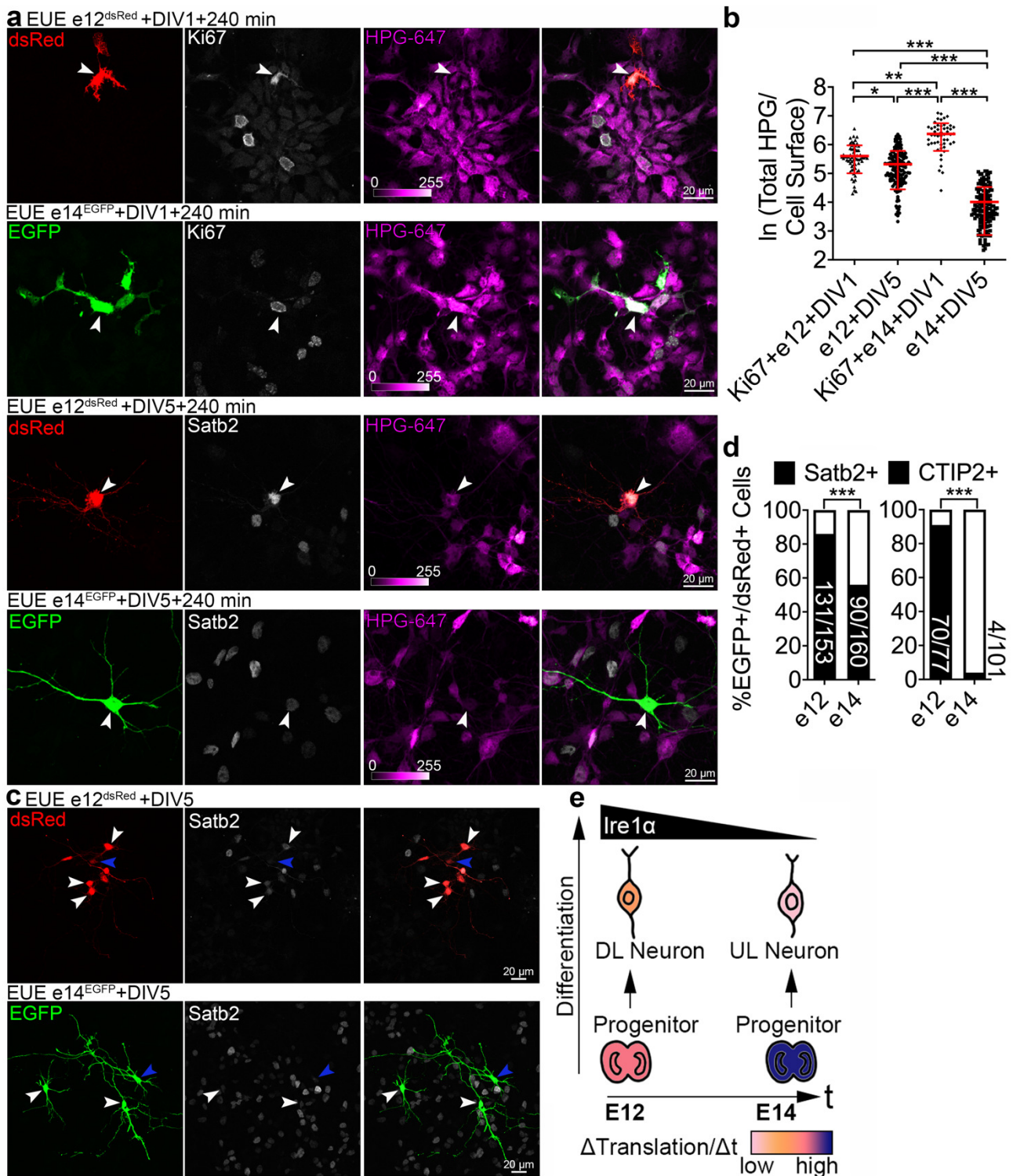


Fig. 6. Progenitors of upper layers display dramatically higher translation rates than progenitors of deeper layers.

485 (a) Images of primary DIV1 and DIV5 cortical cells immunolabeled for EGFP, dsRed, Ki67, and Satb2. To target Ki67-positive neuronal progenitors and their derived progeny, cortices of e12 embryos were *ex utero* electroporated (EUE) to express dsRed, and cortices of e14 embryos to express EGFP. Cells were tritured, mixed, plated together on a glass coverslip and pulsed with HPG for 240 min prior to fixation at DIV1 and DIV5. White arrowheads indicate representative cells. (b) Incorporation of HPG
 490 was quantified as intensity of fluorescence signal normalized to the cell surface area (natural logarithm scale). (c) e12 and e14-derived cells were fixed at DIV5 and immunolabeled for Satb2. White

arrowheads point to neurons expressing high levels of Satb2, blue ones indicate example neurons with lower Satb2 expression. (d) Quantification of the cell identity markers in DIV5 neurons. (e) Schematic summarizing the translation rates ($\Delta\text{Translation}/\Delta t$) in different types of cortical cells for e12- and e14-derived lineage. The triangles indicate the dynamics of developmental enrichment for Ire1 α ; deeper layers (DL) and upper layers (UL). Graph on (b) indicates mean \pm S.D. Statistics for (b), D'Agostino-Pearson normality test and Kruskal-Wallis test with Dunn's multiple comparisons test; (d), Fisher's test. *** $p < 0.001$; $0.001 < ** p < 0.01$; $0.01 < * p < 0.05$.

500 **Transient attenuation of translation results in loss of axon and upper layer type identity**

We then asked if pharmacologically attenuating translation early in development alters the specification of upper layer type neurons. Given a significant difference in translation rates at DIV1 between e12- and e14-derived neuronal cultures, we nucleofected the primary cells to express dsRed or EGFP as described above and exposed them to DMSO or cycloheximide (CHX), an inhibitor of protein synthesis, immediately after plating transiently for 24 hours. Following this, we changed the cell culture medium to one lacking CHX and maintained the primary neuronal cultures until DIV5 (Fig. 7a). Remarkably, transient inhibition of translation affected the appearance of Satb2 expression in cells derived from both stages but had no effect on the expression of CTIP2 (Fig. 7b).

510 Additionally, neurons transiently exposed to CHX failed to project a long neurite at DIV5 (Fig. S5a). While control EGFP-expressing upper layer neurons displayed an asymmetric morphology with a longest neurite enriched for Tau-1 (Fig. S5a), most neurons transiently exposed to CHX failed to break their symmetry and expressed Tau-1 in the perinuclear regions of soma, resembling the scenario upon *Ire1α* (Fig. S3a, bottom panels).
515 We quantified a CHX-induced dose-dependent loss of axons (Fig. S5b) and overall loss of neurite branching (Fig. S5c and S5d).

Altogether, transiently inhibiting protein synthesis at early developmental stages leads to irrevocable loss of Satb2 identity and loss of axons, indicative of critical translational window for both processes and reminiscent of *Ire1α* KO.

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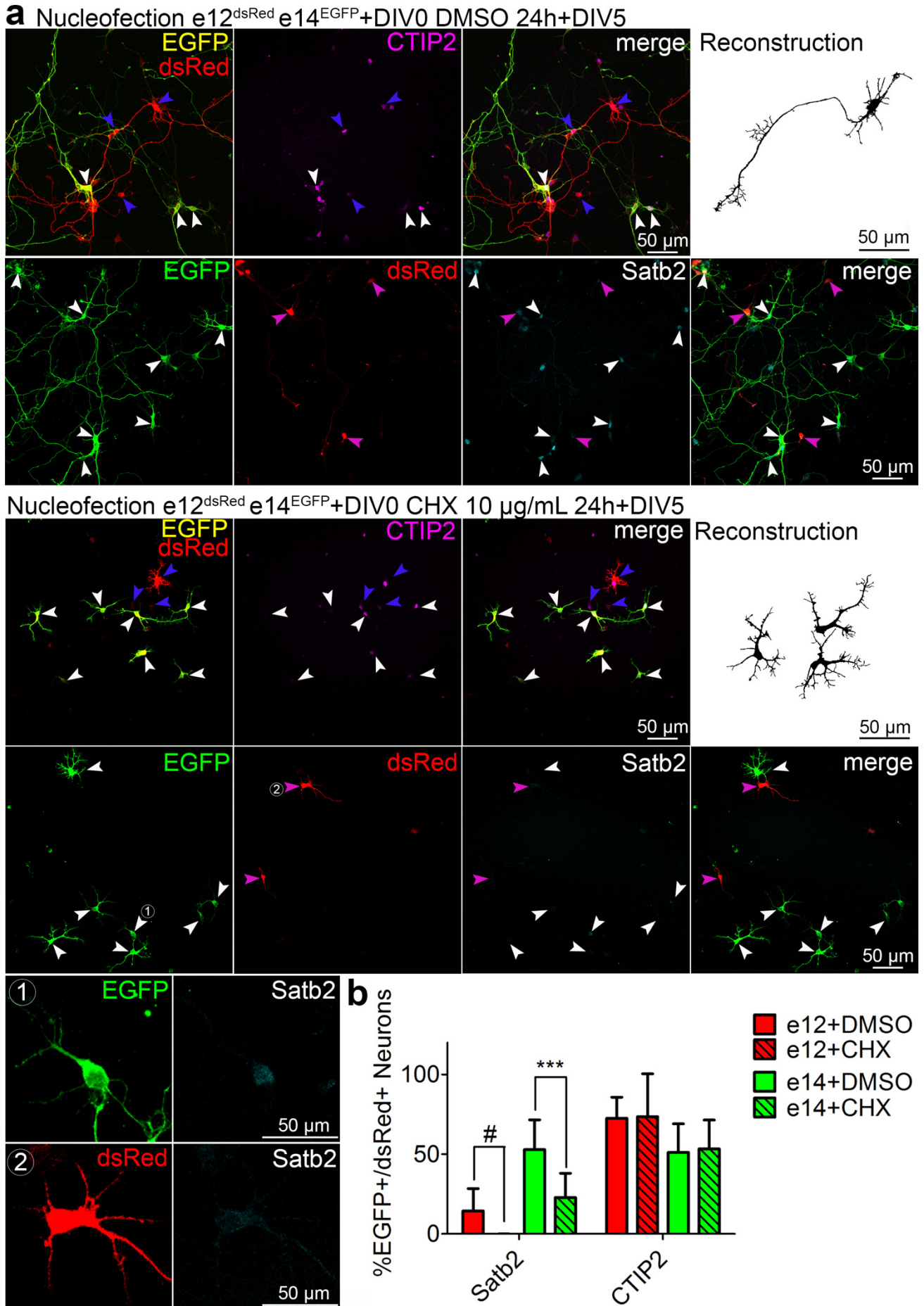
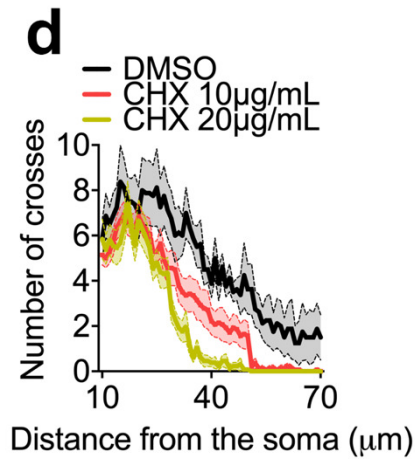
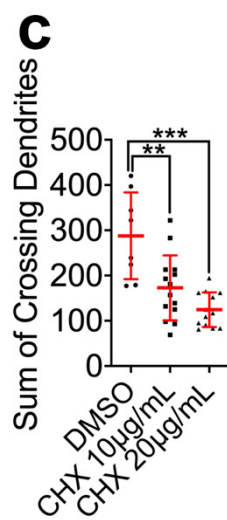
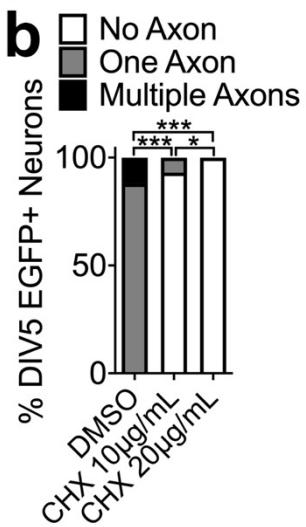
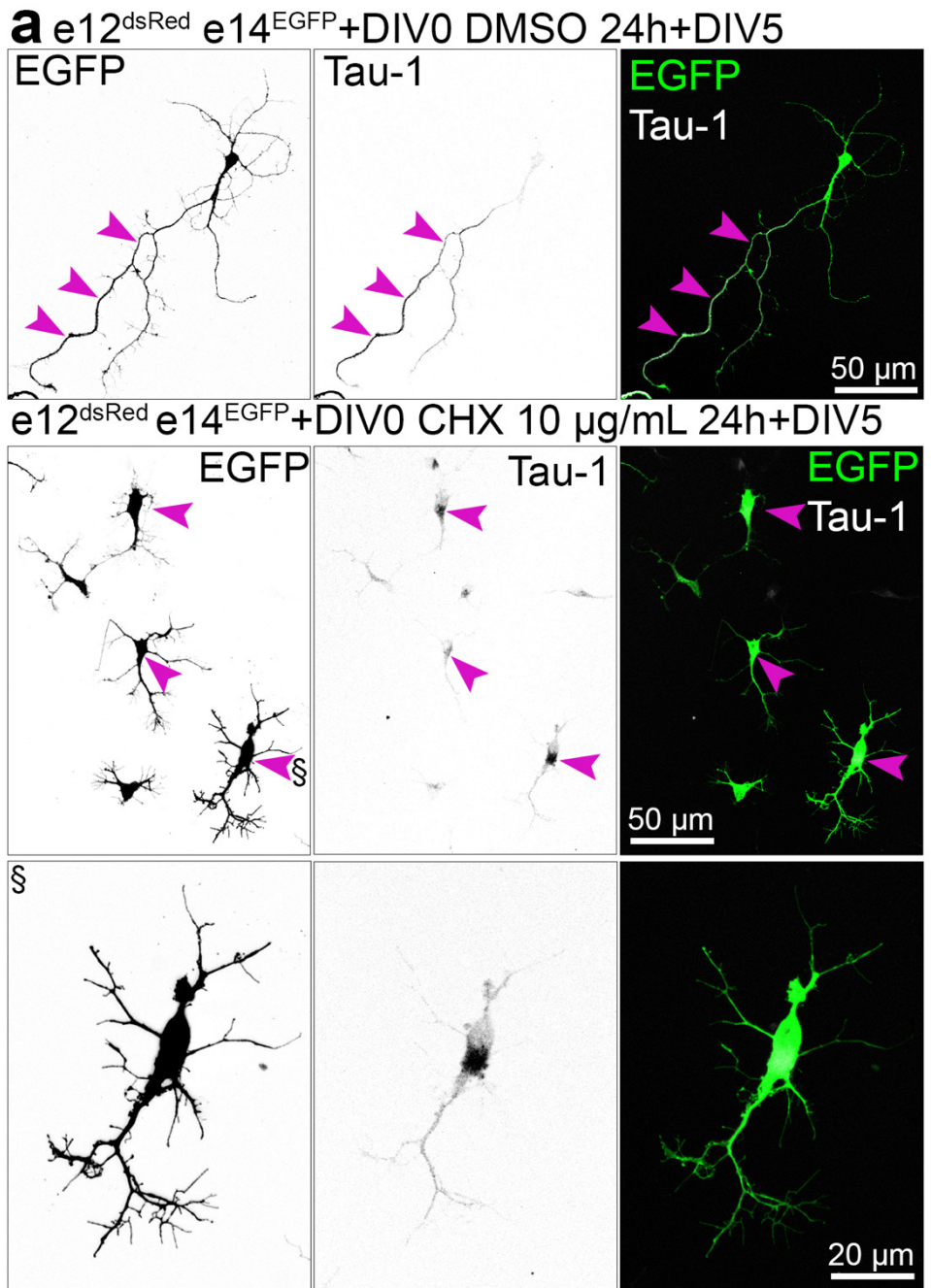


Fig. 7. Upper layer identity requires a critical window of protein translation in precursor cells.

(a) Images of immunolabeled primary cells from e12 embryo nucleofected to express dsRed, and from e14 embryo to express EGFP, mixed and plated on a single glass coverslip. Two hours post-
525 plating, cells were treated with vehicle (DMSO) or cycloheximide (CHX) for 20 hours, followed by medium change, and fixed at DIV5. Upper panels show staining using rat anti-CTIP2, goat anti-EGFP, and rabbit anti-RFP, the latter one recognizing both EGFP and dsRed. For this reason, e12 cells in this case were recognized as solely expressing dsRed (blue arrowheads), but the e14 ones, both EGFP and dsRed (white arrowheads). Lower panels show anti-Satb2, anti-EGFP and anti-dsRed
530 immunostaining with no cross-reacting antibodies; in this case, e12-derived cells express dsRed (blue arrowheads) and e14 ones EGFP (white arrowheads), as expected. Representative neuronal morphology is demonstrated as a semi-automated, EGFP- or dsRed-based reconstruction. (1-2) Example cycloheximide-treated e14 (1) or e12 (2) cortex-derived cells immunolabeled with an antibody anti-Satb2. (b) Quantification of the cell identity markers in DIV5 neurons derived from e12
535 or e14 cortex. Bar graphs indicate mean \pm S.D. For statistics, two-way ANOVA with Bonferroni's multiple comparisons test. # indicates a comparison between fractions of Satb2-positive e12 DMSO- and e12 CHX-treated group, the latter represented by virtually no positive cell. *** $p < 0.001$.

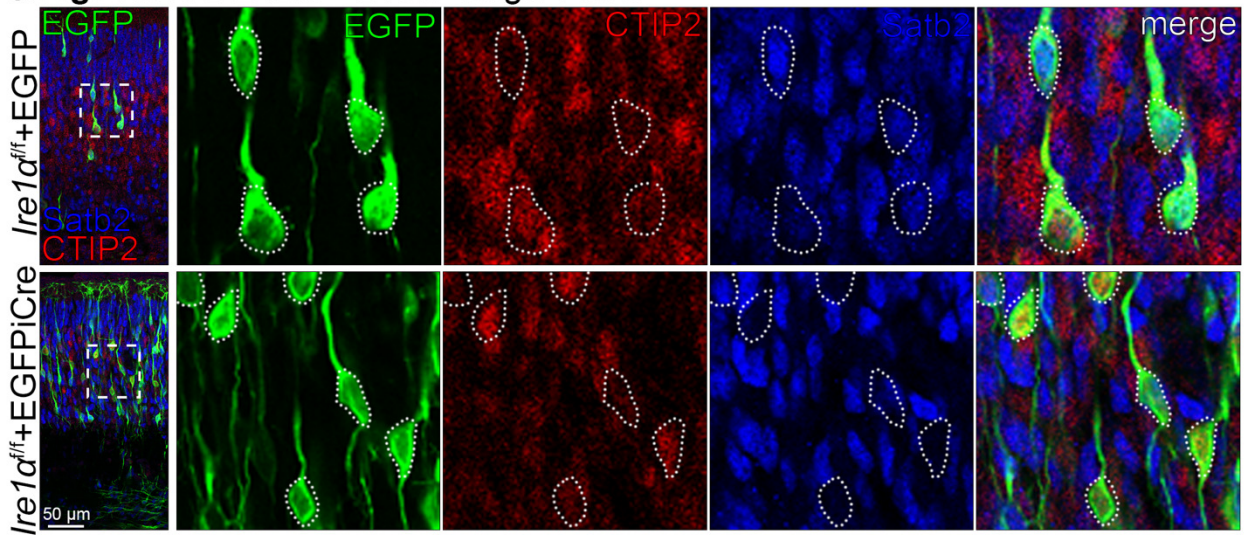


540 **Fig. S5. Related to Fig. 7. Axon formation and dendritic growth requires a critical window of translation in precursor cells.**

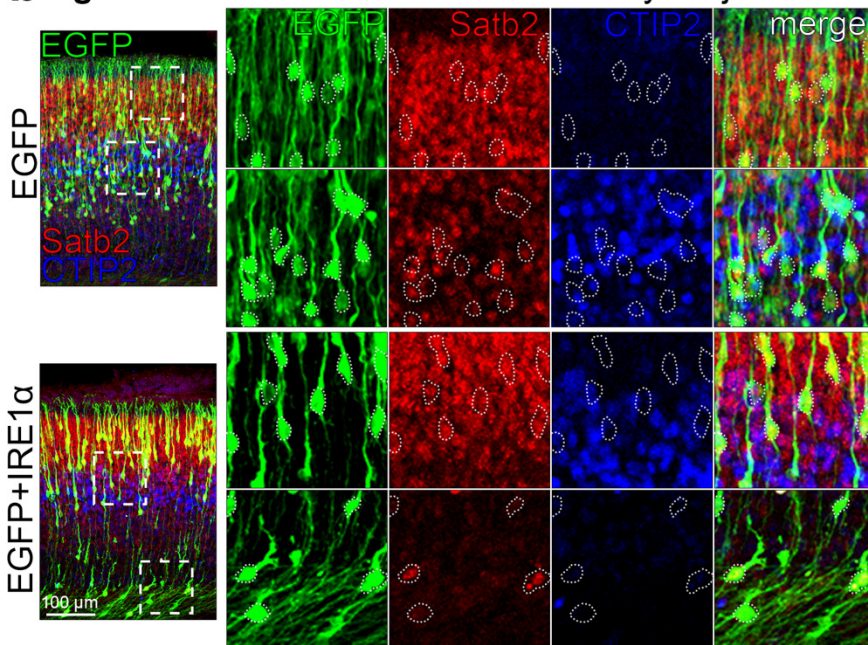
(a) Images of immunolabeled primary cells prepared as described in the legend of Fig. 7a. Cells were fixed at DIV5 and immunolabeled for EGFP, dsRed and Tau-1. Purple arrowheads indicate enrichments of Tau-1. The neuron marked with a paragraph symbol is magnified on the bottom panel.

545 (b) Quantification of neuronal polarity. Axon was defined as the neurite with prominent Tau-1 enrichment [compare with top panel in (a)]. (c-d) Quantification of dendritic complexity in DMSO- and CHX-treated DIV5 neurons. Red line and error bars on (c) indicate mean \pm S.D. Results on (d) are represented as averages \pm S.E.M. For statistical analyses, (b), Chi-square test; (c), D'Agostino and Pearson normality test and one-way ANOVA with Bonferroni post-hoc test. *** $p < 0.001$; $0.001 < **$
550 $p < 0.01$; $0.01 < * p < 0.05$.

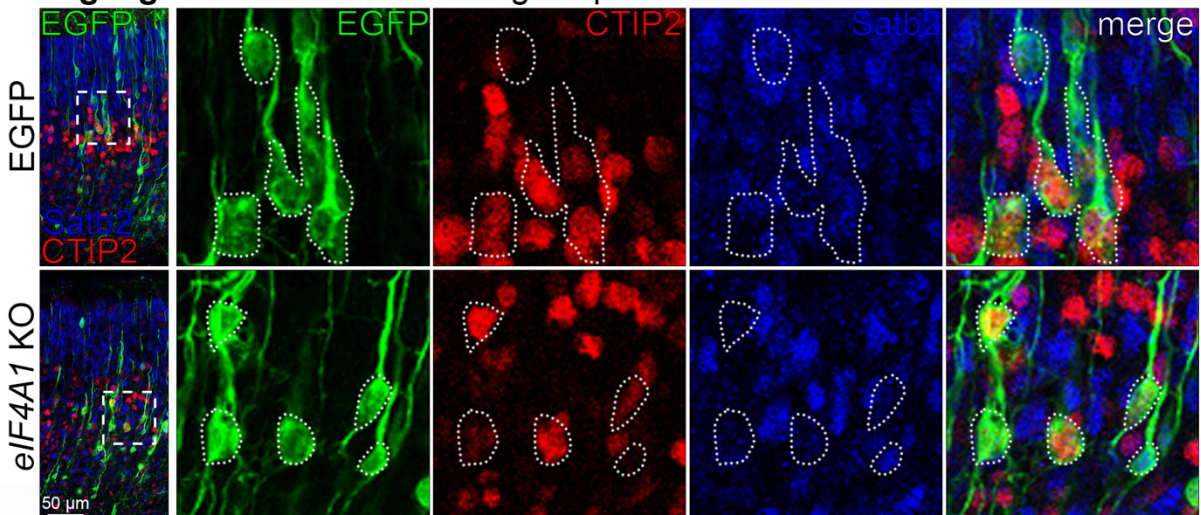
a Fig. 1e. IUE E12.5 - E16.5 Single Plane



b Fig. 1h. IUE E12.5 - E18.5 Max Intensity Projection



c Fig. 3g. IUE E12.5 - E18.5 Single Optical Plane



555 **Fig. S6. Related to Fig. 1, Fig. 3. Zoom-ins for the neuronal identity phenotypes.**

(a-c) Representative images of immunostaining against EGFP, Satb2, and CTIP2 in coronal cortical sections after IUE at E12.5 for indicated conditions. Each panel represents a sample image pair used for identity analysis in experiment indicated on top. The squares depict the area of the zoom-in, outlined are somata of electroporated neurons. Neuronal identity was quantified in the proportion
560 of all electroporated neurons in a given coronal section, here shown are snippets for a visualization of immunostaining.

565 DISCUSSION

Development of the cortex comprises an orchestrated series of sequential, tightly controlled gene expression events with many layers of regulation. Our data implicate the regulation of protein synthesis rate by Ire1 α in the specification of upper layer neurons and their polarization. Through identifying a developmental role for Ire1 α beyond its canonical role in translation stress pathways, we make the surprising finding that different protein synthesis rates are intrinsic features of distinct progenitor and differentiated neuron lineages.

The demand to swiftly and efficiently synthesize and remodel the proteome during cortical development places homeostatic pathways on the brink of cellular stress signaling. We speculate that such demands in normal development are met by the same molecular players as stress pathways, however, likely leading to unique downstream pathways active in development, such as a *Satb2* program. During cortical development, Ire1 α supervises the cellular translation flux (Fig. 4) by driving the expression of translation regulators and RNA binding proteins, essential for the high protein synthesis rate in upper layer progenitors, enabling translation of *Satb2* in their postmitotic progeny (Fig. 3) and proteins crucial for specification of an axon. By labeling the amount of HPG incorporated into the newly synthesized proteins in a given time period, we quantify the total translation flux in a cell. HPG incorporation reflects several translation-regulating processes, e.g. the translation efficiency, polypeptide chain elongation, codon dwell times, overall representing the rate of protein synthesis (Fig. 4).

Seemingly, on top of elegant transcriptional regulation (Nowakowski et al., 2017; Telley et al., 2019), generation of upper layer neurons relies on specific pathways for translational control. One possible scenario involves specific features of 5'UTR regions in genes critical for upper layers (Hinnebusch et al., 2016) and activity of translation initiation complexes, comprising eIF4A, which unwind cap-proximal regions of mRNA prior to its ribosomal loading (Andersen, 2006). In line with this are our observations on decreased translation rates upon *eIF4A1* KO (Fig. 4c and 4d) associated with defects in upper layer neurogenesis (Fig. 3g and 3h).

The *Ire1 α* cKO partially mimics the *Satb2* KO (Britanova et al., 2008), with the loss of upper layer type neurons at the expense of CTIP2-expressing deeper layer cells (Fig. 1 and S2). Strikingly, in the *Ire1 α* cKO cortex CTIP2-expressing neurons are specified during late neurogenesis (Fig. S2e-S2h). Wild type neuronal progenitors at e12.5 generate CTIP2 and *Satb2* positive cells, and as development progresses, late progenitors give rise to *Satb2*-expressing progeny (Fig. S2 and 5-6). Such progressive restriction of neuronal progenitor potency is abolished upon *Ire1 α* loss-of-function. In contrast, high activity of IRE1 α correlates with lower expression of stemness markers, like SOX2, POU3F2 or OLIG2 (Doultsinos et al., 2019).

It is worth emphasizing the results in Fig. 5 and 6 represent different ways of lineage labeling (bulk nucleofection versus clonal progenitor tree), which accounts for different fate signatures in the postmitotic progeny. Our approach to mix labeled cells derived from embryonic cortices of different developmental stages allows for exposure to identical culturing conditions. An interesting future direction is to examine how cellular and environmental composition affects the identity of progenitors, neurons, and associated translation rates.

Global attenuation of translation during the UPR is linked to PERK-mediated phosphorylation of eIF2 α at Ser51 (Krishnamoorthy et al., 2001). The stress response alters the cellular translation machinery and its affinity towards open reading frames of specific type, position, and secondary structures (Starck et al., 2016; Vattem and Wek, 2004). We report that loss of *Ire1 α* results in lower rates of protein synthesis in neuronal progenitors and neurons, independently of eIF2 α , or its canonical downstream splicing client Xbp1 (Fig. S4a and S4b). *Ire1 α* regulates the amount of translating polysomes, with higher levels in the cKO. In the context of lower translation flux, this likely represents ribosome stalling and/or slower elongation, reminiscent of observations in *Huntingtin* mutants (Eshraghi et al., 2021). As *Ire1 α* binds 80S ribosomes directly with high affinity (Acosta-Alvear et al., 2018), and is in the vicinity of the ribosomes embedded in the ER by the translocon, it is well positioned to impact protein synthesis directly during cortex development.

Protein synthesis inhibition with CHX resembles the *Ire1 α* cKO and *eIF4A1* KO (albeit engendering expected stronger phenotypes, given the global inhibition of translation by the compound), with defects in *Satb2* expression and axonal targeting of Tau-1, leading to the formation of symmetrical, non-polarized neurons and ablation of *Satb2* (Fig. 7 and S6). Both the loss of *Ire1 α* and CHX spare the expression and subcellular localization of deep layer neurons-associated CTIP2 and dendrite-enriched MAP2, indicative of the specific requirement for *Ire1 α* and high translation rates during upper layer neurogenesis and polarization. In line with this, we find that late neuronal progenitors display a drastic increase in translation rates.

Apart from being a site of protein synthesis, the ER has been also reported as an organelle crucial for neuronal polarization (Fariás et al., 2019). Upon *Ire1 α* KO, we report a mislocalization of the ER to the polarizing neurite (Fig. 2m and 2n). Defects in axon specification, and thus axonal targeting of Tau-1, AnkG and Navs (Fig. S3) might indicate ER dysfunction upon *Ire1 α* loss, implicating its role in ER integrity and neuronal polarization (Carter et al., 2021). Loss of polarization might also partially explain the filamin A-dependent *Ire1 α* loss-of-function phenotype in laminar positioning of cortical neurons (Urrea et al., 2018). Emergence of either multiple axons or more commonly no axons in *Ire1 α* KO neurons (Fig. 2) implies a disruption of distinct axon polarity-regulating processes in this mutant, likely involving cellular sorting of axonal proteins and ER-dependent microtubule stability, overall leading to disturbed transmission of action potentials. Cortical neuronal migration and neuronal polarization involves *Ire1 α* -driven control of multiple downstream processes, filamin A anchoring on one hand, eEF-2- and eIF4A1-regulated translation on the other hand.

Taken together, our study reinforces the powerful impact of post-transcriptional mechanisms in cortex development. Translational regulation of gene expression during neuronal specification is layered on top of transcriptional mechanisms, many of which have been highlighted in excellent recent work (Nowakowski et al., 2017; Telley et al., 2019). Modulation of translation rates may be a particular requirement for upper layer specification and morphology, evolving as a mechanism to specify upper layers in the late stages of development while restricting the potential of neural progenitors to this lineage. We speculate that a swift upregulation of translation flux in late progenitors might be necessary to efficiently synthesize determinants of upper layer neurons. In the light of eIF4A1 requirement for *Satb2*

identity, it is possible that 5'UTR-embedded structural features are present in mRNAs critical for differentiation of upper layer neurons. Given developmental downregulation of Ire1 α , it is plausible that other translational regulators alongside Ire1 α drive the increased translation efficiency in upper layer progenitors. Thus, translational control by in part by Ire1 α is deployed
655 in normal development to promote upper layer neurogenesis, generate their axons, and wire the mammalian cortex.

SOURCE DATA

Sequencing data are available on NCBI under GSE172489.

660

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AUTHOR CONTRIBUTIONS

670 M.C.A. carried out and conceptualized the study, with significant contribution from E.B. V.T. designed the inhibitor screening, initiated, and supervised the project. A. G. N. performed the experiments with RNAseq and the bioinformatics. M. L. K. purified the ribosomes ex-vivo and analyzed their profiles. T. S. performed the molecular cloning. R. D. significantly contributed to histological and cell culture experiments. M. B. recorded from autaptic cultures and
675 analyzed the results. P. T. shared his expertise and protocols for the FUNCAT and click chemistry. D. K. helped with the initial screening and share the molecule library. O. J. performed proteomic analyses. T. I. shared the *Ire1 α ^{ff}* mouse line. M. R., C. M. T. S., and C. R. supervised works in their labs.

680 CONFLICT OF INTEREST

The authors declare no competing interests.

STAR METHODS

CONTACT FOR REAGENTS AND RESOURCE SHARING

685 Further information and requests for resources and reagents should be directed to Victor Tarabykin (victor.tarabykin@charite.de). Detailed information on the materials used in this study can be found in Key Resource Table.

EXPERIMENTAL MATERIAL AND SUBJECT DETAILS

690 **Animals**

Mouse (*Mus musculus*) lines described in this study have been maintained in the animal facilities of the the Charité University Hospital and Lobachevsky State University. Wild type mice were of NMRI strain. In *Ire1 α ^{ff}* mouse line (Iwawaki et al., 2009), exons 20-21 of *Ire1 α* were engineered to be flanked by loxP sites. To inactivate *Ire1 α* in the developing cortex, we

695 crossed *Ire1a^{ff}* mice with the *Emx1^{Cre/+}* line, in which Cre recombinase is expressed from the
Emx1 gene allele (Gorski et al., 2002). As described in our previous work, for the breeding
we exclusively used *Ire1a^{ff}* males and *Ire1a^{ff}; Emx1^{Cre/+}* females to circumvent leaky
expression of Cre (Luo et al., 2020). Animals homozygous for the loxP alleles in both lines
700 phenotypic changes in the cage environment. For experiments with tdTomato reporter (Fig.
1a-1b), *Satb2^{Cre/+}* (Britanova et al., 2006) males were mated to NMRI wild type females. The
date of vaginal plug was counted as E0.5. All mice were sacrificed by administering a lethal
dose of pentobarbital.

All experiments were performed in compliance with the guidelines for the welfare of
705 experimental animals approved by the State Office for Health and Social Affairs, Council in
Berlin, Landesamt für Gesundheit und Soziales (LaGeSo), permissions G0079/11, G206/16,
and G54/19, and by the Ethical Committee of the Lobachevsky State University of Nizhny
Novgorod.

710 **Sex and age/developmental stage of animals for *in vivo* experiments**

Littermates of both sexes were randomly assigned to experimental groups during
experimental procedures or collection of embryonic tissue. Developmental stages or stages
at experimental interventions are listed on the figures or in the figure legends.

715 **Murine primary neurons**

Embryonic brains were dissected on ice-cold Hank's Balanced Salt Solution (HBSS) under
stereomicroscope to collect cortices in 5 mL of ice-cold 0.25% trypsin (Gibco) solution. After
incubation for 20 min at 37°C with occasional swirling, digestion was terminated by addition
of 1 mL of fetal bovine serum (FBS, Gibco) and 1U of DNaseI (Roche) was added to the tube
720 for 1 min. Next, cortices were carefully washed twice with 5 mL of pre-warmed (37°C)
Complete Neurobasal medium. Further, cortices were carefully triturated in 1 mL Complete
Neurobasal medium 10 times using P200 pipette tip. The debris were then let sink for 1
minute and 150 µL the supernatant transferred to 900 µL of fresh Complete Neurobasal at
37°C. After second round of trituration, cells were counted with Naubauer counting chamber
725 and neurons were seeded in 24 well-plate formats at 120 000 cells (EUE) or 240 000 cells
(after nucleofection) per coverslip in 1mL of Complete Neurobasal medium. The day of
neuronal prep was counted as day *in vitro* 0 (DIV0). Neurons were cultivated at 37°C in the
presence of 5% carbon dioxide in HERA-cell240 (Heraeus) incubator. For the experiments
with DMSO and CHX, neuronal cultures were treated with the substance two hours post-
730 plating. Given substance was thoroughly mixed with fresh medium and placed on attached
cells.

Autaptic hippocampal cultures were prepared from P0 *Ire1a^{ff}* mice as described (Giusti et
al., 2014).

735 **Complete Neurobasal:** 500 mL Neurobasal (Gibco, Life Technologies), 10 mL B-27 (Gibco,
Life Technologies), 5 mL GlutaMAX (Gibco, Life Technologies), 5 mL penicillin/streptomycin
(Life Technologies, Gibco), 1 mL Primocin (Invitrogen).

METHODS

740 **Lentivirus production and infection of autaptic hippocampal cells**

Production of lentiviral particles and infection of autaptic cultures was performed by the Charité Viral Core Facility according to standard published protocols.

Electrophysiological Recordings

745 Voltage and current clamp experiments were performed as described before (Giusti et al., 2014).

Expression vectors

For our screening, we used beta-actin driven expression constructs pCAG-EGFP and pCAG-
750 flox-stop-flox-tdTomato, which are described in our previous work in detail (Ambrozkiwicz et al., 2017)

Cloning strategies for constructs generated in this study

Cloning of all expression vectors in this study was performed using the NEBuilder system
755 according to manufacturer's protocol (New England BioLabs). DNA fragments were amplified using GXL Prime Star DNA polymerase (Takara) using cDNA libraries as templates. Destination vectors were linearized with EcoRI-HF (New England BioLabs).

pRai-HA-IRE1 α

760 Human IRE1 α cDNA (NM_001433.3) was amplified from plasmid template (Addgene, #13009) using the following oligos: 5'- agattacgctatctgtacaggcATGCCGGCCCGGCGGCTG-3' and 5'- ggccgctagcccgggtaccgCTTGTTTGGGAAGCCTGGTCTCCCTGC-3' and inserted into the modified pRaichu vector (Ambrozkiwicz et al., 2020).

765 pCAGIG-6XHis-eEF2

Murine eEF2 cDNA (NM_007907.2) was amplified from cDNA library using the following oligos: 5'-gtctcatcattttggcaaagATGCATCATCATCATCATGTGAACTTCACAGTAGATC-3' and 5'-cggccgcgatattcctcgaggCTACAGTTTGTCCAGGAAGTTG-3' and inserted into pCAGIG vector for simultaneous expression of 6XHis-tagged eEF2 and EGFP.

770

Nucleofection of primary neurons

Transfection of primary cortical cells was performed using nucleofection (Lonza Bioscience) according to our previously published protocol (Ambrozkiwicz et al., 2017).

775 **Small molecule inhibitor screening**

Right after trituration, E13.5 embryonic cortical neurons prepared from *Satb2*^{Cre/+} mice, were nucleofected with pCAG-EGFP and pCAG-loxP-Stop-loxP-tdTomato plasmids and seeded at 120 000 cells per well of 96-well plate. Two hours post-plating, cultures were treated with compounds at two concentrations, in technical duplicates. Cells were then cultivated until
780 DIV2, when the proportion of *Satb2*^{tdTom} neurons normalized to EGFP positive cells was determined using FACS as described (Ambrozkiwicz et al., 2017).

Ex Utero Electroporation (EUE)

Embryonic brains were injected with a glass capillary to fill the entire volume of a single lateral
785 embryonic ventricle with DNA (final concentration of 500 ng/ μ L), followed by electroporation
using 6 pulses of 35V applied using platinum electrodes. The isolated heads were then placed
in ice cold HBSS until proceeding with primary cultures.

Immunocytochemistry

790 Cells cultured on coverslips were fixed with cold (4°C) 4% PFA, 4% sucrose (Merck) in PBS
for 20 minutes in room temperature, washed three times with PBS and incubated with
Blocking buffer for 30 minutes. Next, cells were incubated with primary antibodies diluted with
blocking buffer accordingly for 16-20 hours at 4°C with moderate shaking, followed by
washing 3 times with PBS. Further, secondary antibodies coupled to appropriate fluorophore
795 diluted in blocking buffer were applied for 2 hours at room temperature. After washing 3 times
with PBS, coverslips were briefly rinsed with ddH₂O and mounted on Superfrost Plus glass
slides (Thermo Scientific) with Immu-Mount mounting medium (Shandon, Thermo-Scientific).

Blocking buffer: 5% horse serum (Gibco, Life Technologies), 0.3% Triton X-100 (Roche), in
800 PBS buffer

Fluorescent Noncanonical Amino Acid Tagging (FUNCAT)

For the fluorescent labeling of newly synthesized proteins, we modified previously published
methods (Tom Dieck et al., 2012). In brief, neuronal cultures were fed with 1mM HPG-
805 supplemented Methionine-free Neurobasal Medium, at DIV1 and DIV5, for the time indicated
on the figures and in the figure legends. Cells were then extensively washed in PBS prior to
fixation with 4%PFA/4% sucrose for 20 min. Control and HPG fed neurons were then
fluorescently labelled with 1 μ M Sulfo-Cyanine5 azide (Lumiprobe). To facilitate azide-alkyne
binding we applied sCy5az in a PBS based “click solution” containing 0.2 mM Tris(3-
810 hydroxypropyltriazolylmethyl) amine (THPTA), 20 mM sodium L-ascorbate, and 0.2 mM
copper (II) sulfate pentahydrate (all from Sigma-Aldrich). Cells were incubated at room
temperature for 20 minutes, before PBS washing and immunocytochemistry (described
above).

815 Met-free Neurobasal Medium: 500 mL Met-, Lys-, Arg-free Neurobasal (Gibco, Life
Technologies), 10 mL B-27 (Gibco, Life Technologies), 5 mL GlutaMAX (Gibco, Life
Technologies), supplemented with final 1 mM L-HPG (Jena), 0.8 mM Lys, and 0.4 mM Arg.

Isolation of RNA, cDNA library preparation and RNA sequencing

820 Tools and work surfaces used for RNA work were thoroughly cleaned with 70% ethanol
(Sigma) and rinsed with RNA-Zap (Thermo Fisher) before the procedure. Embryonic brains
at E18.5 were dissected in DEPC-PBS and flash-frozen in liquid nitrogen until the purification
procedure. RNA was isolated using RNeasy columns (RNeasy Mini Kit, Qiagen). Further
steps were performed according to the manufacturer's protocols. RNA was eluted from the
825 silica membranes using 30 μ L of molecular biology-grade (MB)- H₂O. Quality and
concentration of the prepared RNA was determined using an Agilent Bioanalyzer. Single or

paired end TruSeq Stranded total RNA libraries were made with Ribo-Zero Gold rRNA depletion and sequenced using Illumina Nextseq 500/550. Raw reads were aligned to GRCm38.p5 using STAR 2.5.3a (Dobin et al., 2013) with parameters --
830 outFilterMultimapNmax 100 --winAnchormultimapNmax 100 --outSamstrandField intronMotif and the gencode vM16 basic annotation transcriptome. A single count table was generated from the resulting bam files using TETranscripts (Jin et al., 2015). Alignment pipeline was executed using Snakemake (<https://snakemake.github.io/>). Counts were then fit to a quasi-likelihood negative binomial generalized log-linear model and tested for differential
835 expression using EdgeR (Robinson et al., 2010).

Biochemical experiments

Details of the protocols used for Preparation of cortical lysates, Protein concentration measurements, Sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE),
840 Western blotting, Quantification of protein levels are described in our previous works (Ambrozkiwicz et al., 2018, 2020).

Polysome purification

Ribosome fractionation and polysome purification from E18.5 cortex was performed as
845 published protocol (Kraushar et al., 2021). Seven brains (14 neocortex hemispheres) of each genotype were pooled per biological replicate, with analysis in two replicates per condition.

Cryosectioning

For all histological procedures, brain sections were prepared on Leica CM3050S cryostat.
850 Prior to cryosectioning, brains were passed through a sucrose gradient to cryoprotect the tissue. Next, brains were frozen in -38 to -40°C isopentane (Roth). Coronal cryosections of 50 µm thickness were collected in PBS/0.01% sodium azide solution. For *in situ* hybridization 16 µm thickness sections were collected.

855 Fluorescence *in situ* hybridization (FISH)

We used RNAscope Technology to detect mRNA of Ire1α according to the manufacturer's protocols (ACD). Prior to hybridization, embryonic brains at E12.5, E14.5 and E18.5 were collected in DEPC-PBS and incubated in 4%PFA/PBS/DEPC for 16-20 hours at 4°C. Brains were then passed through a series of sucrose solutions (10%-20%-30%/PBS) until they reach
860 osmotic equilibrium, embedded in O. C. T. Compound (Tissue-Tek) in a plastic cryoblock mold and frozen on dry ice. Coronal sections of 16 µm thickness were collected using the cryostat.

Immunohistochemistry

865 Fixed brain sections were washed with PBS three times at room temperature prior to the procedure to remove the sucrose and freezing compound residue. The sections were then incubated with Blocking solution for one hour at room temperature, then with the primary antibody and DAPI diluted in blocking buffer for 16-20 hours at 4°C, washed in PBS three times for 30 minutes and incubated with secondary antibody diluted in the blocking buffer for
870 up to four hours at room temperature. Next, sections were incubated with PBS for 30 minutes

three times and mounted with cover glass (Menzel-Gläser) and Immu-Mount mounting medium (Shandon, Thermo-Scientific).

Blocking solution: 5% horse serum, 0.5% (v/v) Triton X-100, PBS

875

BrdU Injection and Immunohistochemistry

Pregnant females at appropriate stages were intraperitoneally injected with BrdU, as described (Tarabykin et al., 2001). Brains were then isolated at p2 from mice of both genotypes (Control: *Ire1α^{ff}* and cKO: *Ire1α^{ff}; Emx1^{Cre/+}*), fixed in 4% PFA, passed through
880 sucrose gradient and frozen in isopentane bath. Free floating cryosections of 50 μm thickness were placed on the SuperFrost Plus glass slides and let dry overnight at +4°C. Antigen retrieval was performed in boiling citrate-based Antigen Unmasking Solution (Vector, pH 6.0) for seven minutes in a microwave followed by 20 min of cooling on ice. Next, sections on slides were immunolabeled according to the protocol.

885

Expansion Microscopy

For the purpose of expansion of coronal brain sections to visualize morphology of EGFP-expressing neurons, we followed previously published protocol (Asano et al., 2018). For immunohistochemistry, we used anti-EGFP antibody (Rockland) at 1:200 and the secondary
890 Alexa488-coupled one at 1:100. After PBS washes, the sections were incubated with AcX crosslinker (10 mg/mL) in 150 mM NaHCO₃ for 20 hours at room temperature, washed with PBS, incubated for one hour with monomer solution (19% sodium acrylate, 10% acrylamide, 0.1% bis-N',N'-methylene-bisacrylamide, 0.01% 4-hydroxy-TEMPO in PBS) and gel was polymerized by addition of 0.2% APS/0.2% TEMED for two hours at 37°C. The section
895 embedded in the gel was then incubated with a proteinase K in a digestion buffer (50 mM Tris-Cl, 800 mM guanidine-Cl, 2 mM CaCl₂, 0.5% Triton X-100 in pH 8.0) for 20 hours at 37°C with gentle agitation. Next, section was transferred to a Petri dish with ddH₂O, which was changed at least five times for two days. The expansion factor was determined before mounting of the specimen. Sections were then transferred onto the glass-bottom chambers
900 (Ibidi) and imaged using Leics Sp8 confocal.

***In utero* electroporation (IUE)**

Prior to IUE, DNA was diluted in TE buffer and mixed with 0.1% Fast Green FCF (Sigma-Aldrich). Final concentration of DNA used for transfecting cortical progenitors was 200-500
905 ng/μL. The details on the procedure are exactly as we described before (Ambrozkiwicz et al., 2018).

Antibodies

For Western blotting, all antibodies were diluted 1:750, apart from anti-β-actin, which was
910 used at 1:2000, in 5% non-fat dry milk in TBS-T buffer (TBS buffer with 0.1% Tween-20). Antibodies against phospho-proteins and anti-Ire1α were diluted in 3%BSA in TBS-T buffer. Secondary antibodies coupled with HRP were diluted 1:10000 in the same buffer as the primary ones.

For immunocytochemistry, all primary antibodies were diluted 1:300 in the blocking buffer,
915 and the appropriate fluorophore-conjugated secondary antibodies were used at 1:750.
For immunohistochemistry, primary antibodies and the respective fluorophore-linked
secondary ones were diluted 1:500 in the blocking buffer.

Data analysis

920 Details concerning Quantitative Western blotting, Axon counting assays *in vitro*, Neuronal
morphometry, Quantification of distribution of cortical neurons and polarity classification are
described in detailed in our previous publications (Ambrozkiwicz et al., 2018, 2020).

Confocal imaging of immunostaining signals and analysis

925 Images of primary cell cultures and brain sections after immunostaining were acquired using
confocal Leica SL, Leica Sp8 or Zeiss Spinning Disc Microscope.

Colocalization with identity markers

Neuronal identity on brain sections was quantified as fraction of all EGFP-expressing
930 neurons. Identity of primary cells on coverslips was determined as fraction of fluorescent
reporter-expressing cell. Based on the numbers of dsRed- or EGFP-expressing cells, we
either quantified at least 4 fields of view for high-density cultures, or imaged a given number
of cells (indicated on the bar graphs or Table S1) and determined the number of cells
expressing a marker protein for cultures with sparse labeling, i.e. after EUE. Compare Fig.
935 S6.

Laminar positioning (% of CP)

We based our positioning analysis on previous published reports (Oberst et al., 2019). To
determine the position of neurons in the cortical plate, confocal images of EGFP signals were
940 first transformed so that the pia is perpendicularly oriented to the horizontal axis. Next,
positions of neurons were marked in Fiji using the Cell Counter plug-in. Using the y-
coordinate, we then expressed the position of a given cell relative to the size of the CP and
plotted it as % CP, with 0% being the bottom of the cortical plate / the subplate (SP) and a
100% being the pial surface / the marginal zone (MZ). Positions of all cells were then plotted
945 individually on the graph. Only brains with comparable electroporation efficiencies were
analyzed.

Quantification of fluorescence intensity

For quantifications of fluorescent intensity, we defined a region of interest based on the
950 fluorescent marker, i.e. dsRed or EGFP signals. Using Fiji software, the raw integrated
density was then normalized to the cell surface area of the outlined cell compartment. For
our quantifications, we imaged the primary cell cultures as Z-stacks and for the analysis, we
used maximum intensity projections.

955 Gene Set Enrichment Analysis (GSEA)

GSEA was performed on total counts from cKO and control polysome fraction RNAseq in
classic mode, testing set, the difference of classes metric, over 1000 permutations with the

c5.go.mf.v7.2.symbols.gmt (molecular function) annotation. This gave 406/772 gene sets upregulated in cKO polysomes where 158 gene sets were significant at FDR<25%. Detailed 960 results are presented in the Table S2.

Statistical analyses

All statistics were performed using Prism Graph Pad software. Detailed information on statistics can be found in the supplementary material. Description of statistical tests, definition 965 of center, dispersion, precision and definition of significance are also listed in the figure legends. Briefly, distribution of datapoints was determined using D'Agostino and Pearson normality test. For normally distributed data, we used unpaired two-tailed t-test, or one/two-way ANOVA with Bonferroni post hoc test; for data with not normal distribution, we used 970 Mann-Whitney test to compare two groups, or Kruskal-Wallis test with Dunn's Multiple comparison test for comparison between multiple groups, unless specified otherwise. For detailed information on all numerical values, sample size and statistical tests used in all experiments, see Table S1.

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Mouse anti-Tau-1	Millipore	Clone PC16C, MAB3420, RRID:AB_94855
Chicken anti-MAP2	Novus	RRID:AB_350528
Goat anti-GFP	Rockland	600-101-215M, RRID:AB_2612804
Chicken anti-GFP	Abcam	ab13970, RRID:AB_300798
Rabbit anti-Ki67	Gift from Prim Singh	N/A
Rabbit anti-Pax6	Millipore	AB2237, RRID:AB_1587367
Rabbit anti-Tbr1	Abcam	ab31940, RRID:AB_2200219
Rabbit anti-Satb2	Generated in-house	N/A
Mouse anti-Satb2	SCBT	sc-81376, RRID:AB_1129287
Rabbit anti-Cux1 (CDP)	SCBT	M-222; RRID:AB_2261231
Rat anti-CTIP2	Abcam	25B6; RRID:AB_2064130
Rabbit anti-Bcl11b	CST	D6F1, RRID:AB_2797823
Rabbit anti-Ire1 α	Abcam	ab37073; RRID:AB_775780

Rabbit anti-Ire1 α	CST	14C10, RRID:AB_2055392
Rat anti-BrdU	Abcam	ab6326, RRID:AB_305426
Mouse anti-Gapdh	Hytest	5G4-6C5; RRID:AB_1616722
Mouse anti-AnkyrinG	Antibodies Incorporated	75-146; RRID:AB_10673030
Mouse anti-Sodium Channel Pan	Sigma-Aldrich	S8809; RRID:AB_477552
Rabbit anti-RFP	Abcam	ab62341; RRID:AB_945213
Rabbit anti-FoxP2	Abcam	ab16046; RRID:AB_2107107
Rabbit anti-P-S6 Ribosomal protein	CST	2211S; RRID:AB_331679
Rabbit anti-FMRP	CST	4317S; RRID:AB_1903978
Rabbit anti-eIF2 α	CST	9722S; RRID:AB_2230924
Rabbit anti-eIF2 α ^{P^{Ser51}}	CST	3398S; RRID:AB_2096481
Rabbit anti-Xbp1S	CST	12782; RRID:AB_2687943
Mouse anti-Vinculin	SCBT	sc-73614; RRID:AB_1131294
Mouse anti- β -actin	Millipore	MAB1501R; RRID:AB_2223041
Mouse anti- β -tubulin	BioLegend	801201; RRID:AB_2313773
Mouse anti-eEF2 ^{P^{Thr56}}	CST	2331S; RRID:AB_10015204
Mouse anti-eEF2	CST	2332S; RRID:AB_10693546
Rat anti-RFP	Chromotek	5f8-100; RRID:AB_2336064
Donkey IgG anti-Mouse IgG (H+L)-HRPO	Jackson ImmunoResearch	715-035-150; RRID:AB_2340770
Donkey IgG anti-Rabbit IgG (H+L)-HRPO	Jackson ImmunoResearch	711-035-152; RRID:AB_10015282
Donkey anti-goat AlexaFluor-488	Jackson ImmunoResearch	705-546-147; RRID:AB_2340430

Donkey anti-rat AlexaFluor-488	Jackson Immunoresearch	712-545-150; RRID:AB_2340683
Donkey anti-rabbit AlexaFluor-Cy3	Jackson Immunoresearch	711-167-003; RRID:AB_2340606
Donkey anti-rabbit AlexaFluor-Cy3	Jackson Immunoresearch	711-165-152; RRID:AB_2307443
Donkey anti-rat AlexaFluor-Cy3	Jackson Immunoresearch	712-165-153; RRID:AB_2340667
Donkey anti-rat AlexaFluor-647	Jackson Immunoresearch	712-175-153; RRID:AB_2340672
Donkey anti-mouse AlexaFluor-Cy3	Jackson Immunoresearch	715-165-150; RRID:AB_2340813
Donkey anti-rabbit AlexaFluor-647	Jackson Immunoresearch	711-605-152; RRID:AB_2492288
Donkey anti-chicken AlexaFluor-647	Jackson Immunoresearch	703-605-155; RRID:AB_2340379
Draq5	Biostatus Limited	RRID:AB_2314341
Bacterial and Virus Strains		
<i>Escherichia coli</i> TOP10 competent cells	Invitrogen	C404052
pLKO-EGFP Lentivirus	Charité Viral Core Facility	N/A
pLKO-Cre Lentivirus	Charité Viral Core Facility	N/A
FISH Probes		
RNAscope Probe Mm-Ern1	ACD, Bio-technie	438031
Biological Samples		
Chemicals, Peptides, and Recombinant Proteins		
Cycloheximide (CHX)	Merck	01810
Dimethylsulfoxide (DMSO)	Merck	D8418
L-Homopropargylglycine (L-HPG)	Jena Bioscience	CLK-1067
FlashTag-488	ThermoFisher	C34554
Critical Commercial Assays		
RNAscope Multiplex Fluorescent Reagent Kit v2	ACD, Bio-technie	323100
NEBuilder HF DNA Assembly Cloning Kit	New England BioLabs	E5520
Translation Initiation Complex Antibody Sampler Kit	CST	4763
Deposited Data		
Experimental Models: Cell Lines		

HEK293T	Leibniz Institute DSMZ-German Collection of Microorganisms and Cell Cultures	ACC 635; RRID: CVCL_0063
Primary murine cortical cells	This paper	N/A
Experimental Models: Organisms/Strains		
<i>Ire1a</i> ^{fl/fl} mouse line	(Iwawaki et al., 2009)	N/A
<i>Ire1a</i> ^{fl/fl} <i>Emx1</i> ^{Cre/+} mouse line	This paper	N/A
<i>Satb2</i> ^{Cre/+} mouse line	(Britanova et al., 2006)	RRID:MGI:3695609
NMRI wild type mouse strain	Charles Rivers and Janvier Labs	RRID:IMSR_TAC:nmri
Recombinant DNA		
Mouse cortex cDNA library (e12, e14, e16)	This paper	N/A
pCX::myr-Venus	gift from A. Hadjantonakis	N/A
pCAG	gift from Jun-ichi Miyazaki	N/A
pCAGIG		Addgene
pRaichu205X	gift from Michiyuki Matsuda	N/A
pCAG-EGFP	(Lois et al., 2002)	Addgene #14883
pCAG-flox-Stop-flox-TdTomato	gift from B. Eickholt	N/A
pCAG-flox-Stop-flox-EGFP	gift from B. Eickholt	N/A
pCAG-EGFPiCre	gift from R. Hugarir	N/A
pCAG-Cre	gift from S. Parathasarathy	N/A
pcDNA3.EGFP-IRE1 α	(Lipson et al., 2006)	Addgene #13009
pRai-HA-Ire1 α	This paper	N/A
pNeuroD-GFP	gift from G. Fishell	N/A
pNeuroD-Cre	gift from S. Parathasarathy	
pX335-sgRNA-EF(Cas9n)-Eif4a1-R	(Ochiai et al., 2020)	Addgene #122345
pX335-sgRNA-EF(Cas9n)-Eif4a1-L	(Ochiai et al., 2020)	Addgene #122346
pCAGIG-6XHis-eEF2	This paper	N/A
EGFP-Sec61 β	gift from Tom Rapoport	Addgene #15108
pCAG-Intron-dsRed	gift from H. Fuchs	N/A
Software and Algorithms		
Fiji	imagej.net	N/A
GraphPad Prism	GraphPad Software	N/A

Photoshop	Adobe	N/A
Snakemake	https://snakemake.github.io/	N/A
STAR	https://github.com/alexdobin/STAR	N/A
TETranscripts	https://github.com/hammell-laboratory/TEtranscripts	N/A
EdgeR	https://bioconductor.org/packages/release/bioc/html/edgeR.html	N/A
GSEA	https://www.gsea-msigdb.org/gsea/index.jsp	N/A

975 N/A not available

SUPPLEMENTARY TABLES

Table S1. Numerical values, sample size and statistics for experiments in this paper.

Table S2. GSEA report for RNA upregulated in cKO polysomes.

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