Dynamic Axonal Translation in Developing and Mature Visual Circuits

Graphical Abstract

Highlights
- Dynamic translatome of retinal axons in vivo matches changing subcellular function
- Adult CNS axons translate mRNAs for synaptic transmission and axon survival in vivo
- Target mRNAs of key translation regulators show developmental co-regulation
- Axon-specific sequence motifs link alternative splicing to axonal translation

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In Brief
Local mRNA translation in axons of developing and adult CNS neurons in vivo shows dynamic regulation, suggesting functional relevance for neural circuit formation and maintenance.

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Dynamic Axonal Translation in Developing and Mature Visual Circuits

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SUMMARY

Local mRNA translation mediates the adaptive responses of axons to extrinsic signals, but direct evidence that it occurs in mammalian CNS axons in vivo is scant. We developed an axon-TRAP-RiboTag approach in mouse that allows deep-sequencing analysis of ribosome-bound mRNAs in the retinal ganglion cell axons of the developing and adult retinotectal projection in vivo. The embryonic-to-postnatal axonal translatome comprises an evolving subset of enriched genes with axon-specific roles, suggesting distinct steps in axon wiring, such as elongation, pruning, and synaptogenesis. Adult axons, remarkably, have a complex translatome with strong links to axon survival, neurotransmission, and neurodegenerative disease. Translationally co-regulated mRNA subsets share common upstream regulators, and sequence elements generated by alternative splicing promote axonal mRNA translation. Our results indicate that intricate regulation of compartment-specific mRNA translation in mammalian CNS axons supports the formation and maintenance of neural circuits in vivo.

INTRODUCTION

RNA localization and local translation are evolutionarily conserved mechanisms employed by cells to control the precise subcellular positioning of nascent proteins. Neurons are highly compartmentalized cells with functionally distinct cytoplasmic/membrane domains (dendrites, axons, and somas), and emerging evidence indicates that localized mRNA translation supports this subcellular differentiation (Holt and Schuman, 2013; Martin and Ephrussi, 2009). Recent in vitro studies revealed an unexpectedly large population of mRNAs in axons, and inhibiting the translation of just one or two of them can cause specific defects in fundamental axonal behaviors, such as neurotrophin-induced outgrowth, branching, cue-induced chemo-
is supported by mRNAs whose translation is developmentally regulated. We also found that axonal mRNA translation continues in adulthood, when regulators of neurotransmission and axon survival are locally translated. Bioinformatic analysis of key translational regulators, such as mammalian target of rapamycin complex 1 (mTORC1), fragile X mental retardation protein (FMRP), and adenomatous polyposis coli (APC), reveals that their target mRNAs are translationally co-regulated in a stage-specific manner. In addition, axonally translated mRNAs show extensive isoform diversity, yet only one single isoform is usually translated at any given time and these axonally translated isoforms share common regulatory sequence motifs that promote axonal mRNA translation. Collectively, the results provide direct evidence for the occurrence of developmental stage-specific, compartmentalized mRNA translation in developing and mature CNS axons and provide a deeper understanding of the molecular machinery involved in CNS wiring and maintenance.

RESULTS

Retinal RiboTag Labels RGC Axonal Ribosomes In Vivo

In order to isolate mRNAs translated in RGC axon terminals in the SC in vivo, we used the RiboTag knockin mouse line (Sanz et al., 2009), in which Cre-mediated recombination switches the RiboTag allele, which encodes the 60S subunit protein ribosomal protein L22 (RPL22), to the hemagglutinin (HA)-tagged Rpl22 allele (Rpl22-HA). We crossed this mouse with a Pax6-alpha-Cre mouse (Marquardt et al., 2001), which transiently expresses Cre in the neural progenitors in the peripheral retinal primordium, permanently labeling RGCs (Figure 1B; green area in the eye). We confirmed that no resident cells in the SC express Cre by two independent approaches, histological and molecular biological assays (Figures 1C, 1D, and S1; see Supplemental Experimental Procedures). Therefore,
the immunopurification of ribosome-mRNA complexes from the dissected SC allows us to profile local translation in axon terminals of RGCs in vivo (axon-TRAP; Figure 1B).

We sought to visualize the labeled ribosomes using an HA antibody. HA immunoreactivity was observed in the distal neural retina (Figures 1E and S1) and the optic nerve head (ONH) (Figure 1E, white box), the soma-free region where RGC axons collect to exit the eye, indicating that the RGC axons do contain HA-tagged ribosomes. To visualize the tagged ribosomes with higher resolution, we employed immuno-electron microscopy (EM). Immunogold particles specifically labeled a subpopulation of ultrastructurally identifiable ribosomes (Figure 1F) in the distal neural retina in a Cre-dependent manner (Figure 1G). We successfully detected HA-tagged ribosomes in the axon shaft in the ONH and the optic nerve (ON) (Figures 1H and 1I) and presynaptic terminals in the SC (Figures 1J–1L) at all stages, indicating that HA-labeled endogenous ribosomes are transported to the axon, even in the adult.

Together, our histological, molecular biological, and ultrastructural analyses indicate that Retinal RiboTag faithfully labels retinal axonal ribosomes in the SC.

Figure 2. Unbiased Identification of the Axonal Translatome

(A) HA-labeled ribosomes were TRAPed by two independent antibodies against HA, and then co-immunoprecipitated ribosomal proteins from 60S (i.e., rpL24) and 40S (i.e., rpS3a) were visualized by western blot. IgG LC, immunoglobulin G light chain.

(B) Double-strand cDNAs were made from TRAPed RNAs.

(C) Read counts of adult SC samples with or without ribosome run-off. Left panel is a scatter plot of log2 (read count+1), and right panel represents the percentage of genes whose read counts were decreased by run-off.

(D) A scatterplot of log2 (FPKM) between Cre-positive/negative (x axis) and Cre-negative axons (y axis) at stage P0.5.

(E) Change in numbers of DEGs in the retina and axon. For axon, dark pink indicates DEGs at the corresponding stages. Light pink indicates genes that are DEGs only at that stage. Combined value of orange and peach (union of DEGs) indicates the size of axonal translatome.

(F) Somal versus axonal translatomes.

(G) Four different axonal translatomes. See also Figure S2.

An Unbiased Identification of the Axonal Translatome

Because the mRNA bound to the labeled axonal ribosomes of RGCs represents only a small fraction of the mRNA in the SC, a major caveat of axon-TRAP is nonspecific binding of mRNAs derived from the SC cells to immunoglobulins, Protein G, and magnetic beads. To reduce this background noise, we first optimized the immunopurification protocol before performing axon-TRAP. We estimated that approximately 40% of HA-tagged translating ribosomes could be purified with this optimized protocol (Figures 2A, S2A, and S2B; see Supplemental Experimental Procedures). Successful isolation of axonal ribosomes was confirmed by silver staining (Figure S2B) followed by mass spectrometry (unpublished observation), although RPL22-HA pulled down from the SC was below the level of detection by western blot. To assess the levels of background noise, we compared the levels of cDNAs amplified from TRAPed mRNAs (Figure S2C) between
Axon-TRAP Identifies Changing Population of Ribosome-Bound mRNAs in Developing and Mature Axons In Vivo

We used axon-TRAP on SCs dissected out at three specific stages during retinotectal development and in the adult: embryonic day 17.5 (E17.5) (elongating); postnatal day 0.5 (P0.5) (branching); P7.5 (pruning); and adult (mature; Figure 1A). To compare the axonal translatome with the somal translatome, we also analyzed the ribosome-bound mRNAs in dissected Cre-positive retina, which contains the cell bodies of RGCs. When we plotted the normalized read count (fragments per kilobase of transcript per million mapped reads [FPKM]) of each mRNA TRAPed from Cre-positive over Cre-negative SC samples, Cre-dependent signals of mRNAs were immediately visible (Figure 2D, left panel, black dots). This was in contrast with the plot with two biological replicates of Cre-negative SC samples, which showed a clear correlation (Figure 2D, left panel, blue dots). To select Cre-dependent mRNAs in an unbiased way, we performed “differential expression analysis” on biological replicates of Cre-positive and -negative samples using NOIseq, which is well suited for quantitative comparisons for independently performed RNA-seq samples (Tarazona et al., 2011; Figures 2D and S2F; see Supplemental Experimental Procedures). We defined these genes as “differentially expressed genes (DEGs)” (Figures 2D and S2F, right panel, red dots; Table S1) and used these for most of the downstream analyses.

The total number of axonally translated mRNAs was higher in early stages, peaking at P0.5, and decreased postnatally, whereas mRNAs that are translated within the retinal somas showed little change over the periods examined (Figure 2E), consistent with the amounts of axon-TRAPed cDNAs (Figure S2A). Although previous studies demonstrated that proteins are synthesized in developing axons, it has been controversial whether mature CNS axon terminals also have the ability to synthesize proteins at all, partly because of early studies detecting few or no ribosomes in mature axons (references in Piper and Holt, 2004). However, the presence of DEGs, approximately 85% of which were confirmed as being translated (Figures 2C and S2E) and ribosomes (Figures 11 and 1L) in adult axons indicates that axonal mRNA translation persists in adult CNS axons. The axonal translatome of RGCs is largely an evolving subset of the significantly larger somal translatome (Figure 2F), confirming that axon-TRAPed mRNAs originate from RGC neurons. Unlike the somal translatome (Figure S3A), however, the axonal translatome showed extensive developmental regulation (see detailed analysis below) with only 694 out of 2,576 (27%) mRNAs translated at all stages (Figures 2G and S3A), indicating that the axonal translatome is not due to the simple passive diffusion of translated mRNAs from the soma.

Axon-TRAPed mRNAs Encode Axon-Specific Proteome

To discover which classes of mRNAs are preferentially translated in the axon, we performed a gene ontology (GO) enrichment/depletion analysis for genes whose translation level is significantly higher (>100-fold difference) in the axon than in the retina (Figure 3A; “axon-enriched mRNAs”). Reassuringly, analysis with the cellular component category showed that
Axon-enriched mRNAs generally encode proteins that are already known to function in axons, growth cones, and synapses (Figures 3B and S3B). In contrast, mRNAs encoding nuclear proteins (e.g., modifier of chromatin structures) are depleted from axonal translatome. GO terms selectively enriched in the axonal translatome included those involved in vesicle-mediated transport and calcium-mediated signaling (Figure S3C), suggesting that these processes, which play key roles in the distal axon, may be regulated by local mRNA translation.

To explicitly compare axonal and somal translatomes, we used ClueGO software, which reports how many genes in each cluster are assigned with specific GO terms. We compared 2,576 axonally translated mRNAs (“axon-related translatome”) with the same number of mRNAs that are most abundant in the somal translatome but absent in the axonal translatome (“retina-only” translatome). We found that synapse- and axon-related GO terms were generally associated with the axonal translatome, whereas the retina-only translatome was enriched with basal body and nuclear GO terms (Figure 3C). These results indicate the presence of mechanisms for selecting specific mRNAs for axonal translation.

**Axonal Translatome Changes from Axon Elongation to Neurotransmission during Development**

To correlate the local translation with the stage-specific events in axon development, we performed a GO-based analysis for genes that are translated in axons at each developmental stage using 455 neuron-related GO terms (Table S2). The translatome in younger axons (E17.5 and P0.5) was highly enriched with axon-development-related GO terms, including “neuron projection morphogenesis,” whereas that of older axons (adult) was enriched with synaptic-transmission-related GO terms, such as “synaptic transmission” (Figures 4A and S4A; Table S3). The ingenuity pathway analysis (IPA) for canonical signaling pathways also suggests that synapse function was most highly regulated in adult axons (Figure S4B). This result suggests that axonal mRNA translation continues in the mature CNS of mammals in vivo and may regulate presynaptic function.

We found that a number of genes, which are robustly translated in adult axons, encode glutamate receptors and neurotrophin receptors (Figures 4B and S4C), some of which are known to related GO terms, such as “synaptic transmission.”
regulate synaptic transmission in the presynaptic compartment (Pinheiro and Muller, 2008). Furthermore, key components of the trans-SNARE complex, which mediates neurotransmitter exocytosis, are highly translated in mature axons (Figure 4B), suggesting that their local translation plays a role in supporting the core machinery of neurotransmission in presynapses.

Intriguingly, translation of receptors for axon guidance molecules peaks around birth (P0.5) and falls off thereafter (Figures 4B and S4A). Because this is when interstitial branches arise from axon shafts in a topographically biased manner to connect with targets (Figure 1A), stage-specific synthesis of these receptors in the RGC axon may help to fine-tune topographically biased branching. We also noted that the GO terms “neuron remodeling” and “collateral sprouting” were among most enriched in the pruning stage (P7.5). Genes that function in synapse assembly, which include neurexins and presynaptic cell adhesion molecules, were translated in all axons (Figures 4B and S4A).

The Axonal Translatome Changes from Degenerative to Survival Modes at the End of Development

Axon survival is regulated through at least two pathways: by maintaining axonal protein/energy homeostasis and by inhibiting a destruction program mediated by SARM1. Sarm1, which initiates a soma-independent axon destruction program by counteracting Nmnat function (Gerds et al., 2015), is highly translated in developing, but not in adult, axons (Figures 4B and S4C). The same pattern of local translation was observed for caspases, whose local action mediates axon dynamics and developmentally controlled branch destruction (Campbell and Holt, 2003; Campbell and Okamoto, 2013; Simon et al., 2012). These results suggest that developing (arborizing) axons synthesize the components of axon degeneration pathways, perhaps in highly restricted subcellular compartments within the axon, for the selective withdrawal of branches, whereas adult axons shut them off to maintain mature neural connections for long periods of time.

GO terms related to mitochondrial and homeostatic functions, such as “cellular metabolism” and “mitochondrial respiratory chain,” were enriched at all stages, supporting the previous finding that axonal mRNA translation supports mitochondrial function and is required for axon survival (Figures 4A and S4A; Cosker et al., 2016; Hillefors et al., 2007; Yoon et al., 2012). The survival of a neuron whose axon reaches its correct target is regulated by retrograde transmission of a survival signal from the axon terminal, which turns on a transcriptional program for cell survival (Riccio et al., 1997). Previous studies showed that axonal synthesis of transcription factors, such as neurotrophin-induced synthesis of CREB (Cox et al., 2008) and SMAD1/5/8 (Ji and Jaffrey, 2012) and axon-injury-induced synthesis STAT3 (Ben-Yaakov et al., 2012), regulates cell survival during development and in adulthood. Indeed, our IPA analysis revealed that components of these nuclear signaling pathways, including CREB and STAT3 signaling, are enriched in adult axons (Figure S4B). Therefore, our results suggest that local translation promotes survival of mature axons both by supporting mitochondrial function and actively generating survival signals.

Pathological axon degeneration in neurodegenerative diseases has been associated with impaired axonal translation (references in Jung et al., 2012). A KEGG pathway enrichment analysis showed a significant over-representation of genes linked to neurodegenerative diseases, such as Alzheimer’s, Parkinson’s, and Huntington’s diseases (Figure 4A). In particular, we detected robust axonal translation of huntingtin (Htt), Prion protein (Pmp), microtubule-associated tau (Mapt), and amyloid beta precursor protein (App), whose aggregates are strongly associated with neurodegenerative disorders (Figure 4B), suggesting a possible connection between axonal translation and neurodegeneration involving protein aggregation. Intriguingly, activating transcription factor 4 (Atf4), whose excessive axonal translation spreads Alzheimer’s disease pathology across the brain (Baleriola et al., 2014), is also axonally translated at all stages tested. These results support the idea that dysregulated axonal translation may be an underlying cause of neurodegenerative diseases (Jung et al., 2012).

Targets of mTORC1, FMRP, and APC Show Translational Co-regulation in a Stage-Specific Manner

We have shown that the axonal translatome is dynamically regulated during development, and this raises the important question of how axonal translation is controlled by upstream signaling pathways. To investigate this, we performed IPA upstream regulator analysis, which is based on published data of gene knockdown or knockout studies where protein products were measured when translational regulator function was impaired. mTORC1 activity was predicted to peak in actively wiring axons, as its target mRNAs showed a steep increase at P0.5 (Figure S5A), consistent with previous studies demonstrating that axonal mRNA translation is regulated by mTORC1 (Campbell and Holt, 2001) and required for axon branching (Spillane et al., 2013). In contrast, the activity of FMRP was predicted to peak later at P7.5, because its target mRNAs (whose translation is repressed) showed a coordinate decrease in translation in mature axons (Figures 5A, 5B, S5A, and S5B). This result suggests that the translational brake mediated by FMRP is utilized in maturing CNS axons as in dendrites (Bagni and Greenough, 2005; Darrell and Klann, 2013). Consistent with this result, known targets of FMRP and mTORC1 in the axonal translatome showed clearly different translational patterns from the non-target mRNAs: their translation increased at P0.5 (Figure 5A, left panel, red and blue; median shifts right) and decreased at P7.5 (right panel). Another intriguing translational regulator was APC, which was recently shown to regulate microtubule assembly and axonal growth by local translation (Preitner et al., 2014). Our analysis indicates that the translation of APC target mRNAs is highest in the youngest axons (E17.5) and steadily decreases thereafter (Figures 5A and 5B), consistent with the primary role of microtubule assembly in axon growth. In contrast, the targets of TDP-43 and FUS, well-known neuronal RNA-binding proteins (RBPs), showed a distribution not significantly different from the total axonal translatome (Figures 5B and S5C), although the possibility remains that TDP-43 and FUS regulate stage-independent axonal mRNA translation. A principal-component analysis (PCA) also showed clear separation of the mTORC1, FMRP, and APC targets from the rest of the axonal translatome (Figure S5D). MicroRNAs (miRNAs) make up another class of translational regulators that function in the axon (Sasaki et al., 2014). We found that the translation of miR-1 target mRNAs
decreases as the axon matures, suggesting that miR-1 abundance and/or activity increases during RGC axonal development (Figure S5A).

We took an independent approach to investigate the possibility of developmental-stage-dependent regulation of mTOR and FMRP signaling in RGC axons. We measured the abundance of phosphorylated mTOR (p-mTOR) and S6 (p-S6) in cultured primary mouse RGC axons by quantitative immunofluorescence (QIF), which positively correlate with mTORC1 activity (Copp et al., 2009; Laplante and Sabatini, 2012; Figure 5C). We found that they increased between E17.5 and P0.5, supporting our hypothesis that mTORC1 activity rises in RGC axons during this period. In contrast, the level of FMRP decreased in the same period, in accordance with our model that the translational brake is weakened in P0.5 axons (Figure 5C).

To gain more insight into mRNA-specific translation in the axon, we compared the RGC axon transcriptome of E17.5 (Zivraj et al., 2010) to the axon translatome at the same stage. We analyzed the genes that are detected in the transcriptome, but not in the translatome, because this group may contain candidates for translationally repressed (“masked”) mRNAs. We found that a significant portion of these candidates was translated at the three later stages tested (P0.5, P7.5, or adult) because their levels in the transcriptome correlated with the probability for translation at later stages (Figure 5D). This suggests the possibility that the mRNAs, which are present in high abundance, but not translated, are being stored for translation in later stages. In contrast, the genes that are present both in the young transcriptome and translatome did not show this trend (Figure S5E).

Strikingly, mRNAs that are “unmasked” at the same stage encode various components of specific signaling pathways (Figure 5D). For example, components of dopamine receptor signaling, WNT/β-catenin signaling, and the oleic acid biosynthesis pathway were specifically unmasked in P0.5, P7.5, and...
Intriguingly, the axonal translatome showed more-extensive diversity of mRNA isoforms than the somal translatome (Figure 6A). To address the possibility of isoform-specific axonal translation, we selected 164 alternative events that produce two isoforms both in the axonal and retinal translatomes. Then, we calculated the “percentage spliced in” (PSI or $\Psi$) values, which represent the fraction of the longer isoform (Katz et al., 2010). $\Psi_{\text{retina}}$ was uniformly distributed ($0 < \Psi < 1$), indicating that there is no clear bias in translational efficacy (Figures 6B and 6C). However, $\Psi_{\text{axon}}$ was biased to the two extremes (i.e., $\Psi = 0$ or $\Psi = 1$), indicating that only one of the two isoforms is selectively translated in the axon (Figures 6B and 6C). Notable examples are Acot7, an acyl-coenzyme A (CoA) thioesterase gene required for lipid biosynthesis and neuron survival (Ellis et al., 2013); Syntaxin 3 (Stx3), a SNARE component gene; and Clta, a clathrin light chain A gene, which show clear axon-specific usage of first, last, and internal exons, respectively (Figures 6D, 6E, and S6A). Intriguingly, axon-specific isoforms of Acot7 and Stx3 encode proteins with slightly different amino acids at the N and C termini, respectively (see gene models in Figures 6D and 6E), suggesting that alternative splicing may couple axon-specific protein isoforms with a unique sequence tag in the UTR.

Unexpectedly, we detected a number of back splicing events for three genes (Rhobtb3, Ubn2, and Ankr12), which indicate the potential presence of circRNAs in the axonal, but not in the retinal, translatome, and we could detect these mRNAs by RT-PCR of unamplified axonal translatome (Figure S6B). Although previous studies suggested that the circRNAs are not translated (Guo et al., 2014), our result raises the possibility that the ribosomes can associate with circRNAs in axons. However, further studies are needed to address whether proteins are actually synthesized from these circRNAs.

**Cis-Regulatory Elements Couple Alternative Splicing with Axonal Translation**

The dominance of a single alternative exon in axons suggests that axonal mRNA translation might be mechanistically linked to alternative splicing. We focused on the axonally enriched mRNAs with an alternative first or last exon because 5’ and 3’ UTRs generally contain localization signals (references in Jung et al., 2012; Figures 6D and 6E). In order to investigate whether the axon-specific exons are sufficient to promote axonal mRNA transport and translation, we used a diffusion-limited, membrane-targeted EGFP (myr-d2EGFP), which is a faithful reporter of local protein synthesis in dendrites (Aakalu et al., 2001) and in axons (Andreassi et al., 2010; Cox et al., 2008). We fused the axon-specific or axon-absent (retina-restricted) alternative exon of each gene to myr-d2EGFP so that a reporter mRNA containing each motif would be generated in cells (Figures 7A and 7B). To test these reporters in RGCs, the same cell type from which they were identified, we used Xenopus primary retinal cultures (Campbell and Holt, 2001), which are amenable to screening multiple motifs. We confirmed that alternative usages of the 5’ and 3’ UTRs of Acot7 and Stx3, respectively, are conserved between mouse and Xenopus (Xenbase and UCSC genome browser). Fluorescence recovery after photobleaching (FRAP) was monitored in the growth cones of cultured RGCs at 1-min intervals for 10 min. Remarkably, the axon-specific
isoforms showed rapid and robust FRAP signals, whereas the retina-specific isoforms did not (Figures 7A and 7B). These results indicate that axon-specific exons of Acot7 and Stx3 are sufficient to promote axonal mRNA translation (Figures 7A and 7B).

We next investigated whether axon-specific exons might contain “generalizable” motifs responsible for axonal mRNA translation. We searched for common sequence elements that are enriched in axon-specific alternative exons (Figure 7C) and in the 5' and 3' UTRs in constitutive exons (Figures S7A and S7B) of axon-enriched mRNAs (Figure 3A). To understand the potential function of identified sequence elements, we searched for genes that contain these elements in the entire mouse genome. Remarkably, the element-containing genes generally encode regulators of axon and synapse function (Figures 7C and S7C). Strikingly, five of six motifs identified from alternative exons and five of twelve motifs in constitutive exons of axon-enriched mRNAs showed a significant FRAP signal at 10 min, indicative of increased axonal mRNA translation of a reporter mRNA when incorporated in the 5' or 3' UTR as in Figure 7B (Figures 7C and S7C). These results suggest the potential links between the sequence elements and axonal mRNA translation and thus provide further insight into the mechanisms underlying the selective and dynamic nature of the axonal mRNA translation.

DISCUSSION

Here, we developed a mouse model of axon-TRAP to isolate mRNAs translated in the distal axon of RGCs in vivo and performed a genome-wide survey of the axonal translatome at critical time points during the assembly of visual circuitry and in adulthood. The axonal translatome is generally a subpopulation of its somal counterpart but is enriched in genes with axon-specific roles. We found that broadly two classes of local translatomes exist in the distal axon: one being constitutively translated and the other being developmentally regulated. The former generally encodes the regulators of protein and energy homeostasis, and the latter encodes proteins required for stage-specific events, such as axon elongation, axon branching, synapse

Figure 7. Cis-Regulatory Elements Link Alternative Splicing to Axonal Translation

(A and B) Axon- and retina-specific Acot7 and Stx3 UTR isoforms fused with myr-d2EGFP were expressed in cultured RGCs (Xenopus). Quantification of fluorescence intensity after photobleaching (FRAP) revealed axon-specific isoforms of Acot7 (A) and Stx3 (B) markedly increase axonal translation of the myr-d2EGFP reporter construct compared to retina-specific UTR counterparts. Data at each 1 min time point represent the mean fraction of recovery relative to pre- and post-bleach levels ± SEM (n = 9 and 10 for axon and eye-specific 5' UTR of Acot7, respectively; n = 14 and 14 for axon and eye-specific 3' UTR of Stx3, respectively). ***p < 0.0001; two-way ANOVA. FRAP signal recovery was abolished by 40 μM anisomycin (10 min post-photobleach: Acot7 axon-isoform + anisomycin 0.064 ± 0.028; Stx3 axon-isoform + anisomycin 0.085 ± 0.026). Representative images of RGC axonal growth cones showing fluorescent recovery after photobleaching for each reporter construct are shown (right). The scale bars represent 10 μm. (C) GO enrichment analysis for entire genome containing axon-specific sequence motifs associated with alternative exons (5' G or C) and their relative efficiency in axonal mRNA translation using myr-d2EGFP reporter constructs. Significance of FRAP recovery curves were compared to no UTR control across 10 min (n ≥ 10 for each construct). Statistical significance of FRAP compared to the no-UTR control was tested across all time points (1–10 min) using a two-way ANOVA (**p < 0.0001 compared to no-UTR control). For representative purposes, the mean fluorescence recovery at 10 min post-photobleaching is shown. Error bars represent SEM. See also Figure S7.
formation, and synaptic transmission. The adult axonal translatome is unique, and its main role is likely to regulate synapse function. Developmentally regulated translatomes were subdivided according to the changes in translation between stages, and those that showed a coordinate change were found to share common upstream regulators, such as mTORC1, FMRP, and APC, as well as novel sequence elements that possibly regulate axonal mRNA translation. Additionally, we found that axonally translated mRNAs were frequently specific splice variants that carried axon-specific motifs. Together, our results show that extensive local mRNA translation occurs in the developing and mature mammalian CNS axons in vivo and provide strong evidence that highly regulated axonal mRNA translation might be at the heart of CNS development and the maintenance of synaptic function.

Previous studies using cultured neurons have revealed that some mRNAs are stored in a translationally repressed state (Buxbaum et al., 2014; Graber et al., 2013). Two independent and complementary approaches have been developed to ask which mRNAs are translated in the axon (Kim and Jung, 2015): metabolic labeling of newly synthesized proteins and isolation of ribosome-bound mRNAs. Proteomic approaches provide the ultimate readout of gene expression, as they can identify post-translationally modified protein products, but a critical limitation of proteomics is that the probe-tagged amino acid or its analogs label all cells, limiting its use to compartmentalized axon culture. An alternative strategy to identify newly synthesized proteins is to look at translated mRNAs (translatome), because these are the obligate precursor to the de novo proteome. A key advantage of this approach is that ribosomes can be isolated from a specific cell type by expressing a genetically encoded epitope-tagged ribosomal protein in just the cells of interest (TRAP; Heiman et al., 2008). Isolated ribosome-mRNA complexes either can be partially digested by RNase, and the fragments protected by ribosome binding can be sequenced (“ribosome profiling”; Ingolia et al., 2011), or the entire mRNA can be separated from the ribosome and directly sequenced. Only the latter method, which we use in this study, allows the discovery of novel isoforms outside the protein-coding region.

We compared translatomes of RGC axons and retinal cell bodies in the same animals. It should be noted that the retinal translatome includes the translatome of the short axons and dendrites of the intraretinal circuitry, as well as their cell bodies. Therefore, the number of mRNAs that are identified as selectively translated in the axon in this study may be an underestimation. Additionally, the expression of tagged ribosomes in non-RGC retinal neurons can potentially introduce bias into the axon/retina translatome. Therefore, the number of mRNAs that are identified as selectively translated in the axon may be an underestimation.

In this study, we show that the RGC axonal translatome changes in a developmental-stage-specific manner, in such a way that proteins playing a key role at specific periods are synthesized when needed. This result is in agreement with a recent study using the Drosophila visual system, which reported that neuronal differentiation associated with maturational presynaptic terminals is regulated by coordinate control of mRNA translation (Zhang et al., 2016), although the subcellular location of mRNA translation was not addressed in that study. Whether signals that regulate mRNA-specific translation come from a cell-intrinsic timer or cell-extrinsic cues remains to be investigated, but our bioinformatic and experimental analyses suggest that this involves stage-dependent activation of RNA-binding proteins, including FMRP. Because FMRP is known to inhibit translation of proteins required for synapse formation and its loss of function leads to over-branching of CNS axons (references in Darnell and Richter, 2012), it is reasonable to assume that FMRP may be activated after CNS axons make appropriate synapses to limit the number of synapses that a single axon makes. In this sense, it is intriguing that defective translational machinery, which is expected to affect all cells in the organism, leads to enigmatically synapse-specific phenotypes, ranging from defective synaptic transmission to impaired cognitive function and memory (references in Buffington et al., 2014). Although the subcellular location of this pathogenesis is unknown, it will be interesting to test whether the axonal translatome of developing cortical neuronal axons in these mouse models of neurodevelopmental disorders is any different from normal mice.

The local mRNA translation in axons of mature neurons has been a subject of long-standing debate (Piper and Holt, 2004). Evidence indicates that ribosomes exist in mature CNS axons (Koenig et al., 2000; Kun et al., 2007; Walker et al., 2012) and that their number is dynamically regulated under normal and pathological conditions (Verheijen et al., 2014). However, what proteins are locally translated in the mature axons was unknown. Our comparative analysis of mature and developing retinal axonal translatomes suggests that local protein synthesis regulates synaptic transmission and axon maintenance. Because axonal translation has been implicated in axonal survival and degeneration (Jung et al., 2012), it will be important to find out whether pathological axon degeneration is preceded by defective axonal translation. The power of axon-TRAP is that it can be extended to other neurons whose cell bodies and axons are anatomically separated. One such example is cortical and spinal motor neurons, whose axonal degeneration leads to human diseases, such as amyotrophic lateral sclerosis (ALS). Recent evidence suggests that defective axonal mRNA transport and translation may be an underlying cause of ALS pathology (Alami et al., 2014; Murakami et al., 2015). Our new technical approach and datasets should provide a valuable resource for future studies.

**EXPERIMENTAL PROCEDURES**

**Histological Analysis**

For immunohistochemistry, tissue sections (12 μm) were visualized using an anti-HA antibody (Abcam ab9110) and a secondary antibody conjugated to Alexa 488 (Life Technologies). For immuno-gold EM, tissues were fixed in 4% paraformaldehyde in 0.1 M HEPES (pH 7.4), and the HA-tagged ribosomes were visualized by the same anti-HA antibody and immunoglobulin G (IgG) conjugated with gold (10–15 nm). Mouse RGC axon culture and QIF were
performed as previously described (Zivraj et al., 2010) using the following antibodies: anti-mTOR (phospho S2448) antibody (Abcam 109268), anti-FMRP antibody (Abcam 17722), and anti-RPS6 (phospho S235 + S236) antibody (Abcam 12864).

Axon-TRAP
A homogenized RiboTag female mouse was mated with a Pax6-alpha-Cre male to produce Cre-positive and Cre-negative mice in a single litter. Three eyes or six SCs were homogenized, and post-mitochondrial fractions were collected. The mRNA-ribosome complexes were precipitated using the polyclonal HA antibody and Dynabeads Protein G (Life Technologies 10004D). For the in vitro ribosome run-off experiments, TRAP was performed after lysis was incubated with rabbit reticulocyte lysate (Promega), harringtonine (Sigma), and 4E1RCat (Sigma) at 37°C for 30 min. Ribosome-bound mRNAs were amplified by a method developed by Tang et al. (2009) with slight modification and sequenced using Illumina HiSeq2000 or NextSeq500. All experiments complied with protocols approved by the University of Cambridge and the Yonsei University College of Medicine Institutional Animal Care and Use Committees.

Data Analysis
The sequence reads were mapped using TopHat 2 version 2.0.12, and FPKM values were estimated using Cufflinks. Read counts for each gene were determined using HTSeq version 0.6.1p1. For the identification of translated mRNAs in RGC axons, we applied differential gene expression analysis on read count using NOISeq. De novo motif analysis was performed using HOMER version 3.0 with custom FASTA files.

A detailed description of all experimental procedures is provided in the Supplemental Information.

ACCESSION NUMBERS
The accession number for the RNA-seq data reported in this paper is GEO: GSE79352.

SUPPLEMENTAL INFORMATION
Supplemental Information includes Supplemental Experimental Procedures, seven figures, and three tables and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2016.05.029.

AUTHOR CONTRIBUTIONS
H.J. and C.E.H. conceived and supervised the project. H.J. performed histological experiments. H.J. and T.S. performed biochemical and molecular biological experiments. J.T.S. performed bioinformatic analyses. J.J., performed Cre specificity experiments, J.O. performed motif imaging and QIF, and B.T.-B. and J.Q.L. performed FRAP and QIF. P.S.A. provided RiboTag mice and the original TRAP protocol. H.J., T.S., and C.E.H. wrote the manuscript.

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We apologize to authors of key papers, which we could not cite due to space limitations. We thank W. Harris for many valuable discussions and critical reading of the manuscript. We thank A. Riccio for myr-d2EGFP; J. Skepper for EM; Gihoon Son for advice on IPA; M. Minett for advice on mouse genetics; and K. Mooslehner, A. Dwivedy, and Jeeun Song for technical assistance. This work was supported by Wellcome Trust Programme Grant (085314/Z/08/Z), European Research Council Advanced Grant (332817) to C.E.H., Cambridge Wellcome Trust PhD programme in Developmental Biology (to B.T.-B.), Gates Cambridge Scholarship (to J.Q.L.), Basic Science Research Program (NRF-2013R1A1A0099625 and NRF-2014K2A7A1036305), Biomedical Technology Development Program (NRF-2013M3A9D5072551), Brain Research Program (NRF-2015M3C7A1028396) funded through the NRF by the Korean government (MSIP), Yonsei University Future-Leading Research Initiative of 2015 (2015-20-0095), and a faculty research grant from Yonsei University College of Medicine for 2013 (6-2013-0064-2-1) to H.J.

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Supplemental Figures

A Cre activity reporter mice

alpha-Cre / Rosa26-LacZ
(labels cell body only)

alpha-Cre / Rosa26-TauLacZ
(labels cell body & axon)

Retina

Optic chiasm (OC)

Superior colliculus (SC)

B Labeled cells and axons in Retinal RiboTag

<table>
<thead>
<tr>
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<th>Rosa26-StopLox-LacZ</th>
<th>Rosa26-StopLox-TauLacZ</th>
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<tr>
<td>Retina</td>
<td>a1, a2, a3</td>
<td>b1, b2, b8</td>
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<tr>
<td>OC</td>
<td>a4, a5, a6</td>
<td>b4, b5, b6</td>
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<tr>
<td>SC</td>
<td>a7, a8, a9</td>
<td>b7, b8, b9</td>
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Figure S1. Specific Labeling of Retinal Axons in Retinal RiboTag, Related to Figure 1

(A) Two Cre activity reporter mice were used in this study. The LacZ reporter labels the cell bodies of Cre-positive cells and their progeny, whereas the TauLacZ reporter labels both the cell bodies and axons.

(B) X-gal staining of the retina, optic chiasm (OC) and superior colliculus (SC) in alpha-Cre; LacZ reporter gene double positive mice. Cre labels most peripheral neural retinal cells in both mice. No cells in the SC used for TRAP in this study express Cre as evidenced by the lack of X-gal stain in the LacZ reporter SC. Unlike the alpha-Cre; LacZ mice, alpha-Cre; TauLacZ mice show robust staining not only in the cell bodies but also the OC and the SC. The SC, which was used for axon-TRAP, is highly innervated by retinal axons.
Figure S2. Axon-TRAP, Related to Figure 2
(A) Bioanalyzer analysis of axon-TRAPed mRNA. Lower tables show the amounts of total RNAs and amplified cDNAs for each TRAPed sample.
(B) Silver staining of axon-TRAPed protein complexes following SDS-PAGE.
(C) Strategy for cDNA synthesis and amplification adapted from the study by Tang et al.
(D) Ribosome run-off experiment. The amplified cDNAs from TRAPed mRNAs with or without run-off (P0.5 retina).
(E) Retinal and axon-TRAP combined with ribosome run-off.
(F) Scatterplots of log₂ (FPKM) between Cre-positive (x axis) and Cre-negative axons (y axis).
Figure S3. Comparison between the Axonal and the Retinal Translatome, Related to Figure 3

(A) The upper panel shows a heat map of hierarchical clustering on the normalized level of axonal and retinal translation of genes. Each row in the heat map corresponds to a single gene. The color of the heat map represents the log2 (FPKM value) for each gene (gray = not detected). The lower panel shows a heat map of a correlation matrix.

(B) Tables showing the ranking of most significantly enriched GO terms in axon-enriched mRNAs and retina-enriched mRNAs. Terms presented in Figure 3B are shown in red.

(C) A heat map showing the enrichment of GO terms in the biological process (BP) category. The colors of the heat map represent the log2 value of the fold enrichment for each GO term value (red = enriched, blue = depleted, gray = not detected), and the numbers on the heat map are $-\log_{10}$ (Fisher’s exact p value) for enrichment.
Figure S4. Gene Set Enrichment Analysis Describing the Developmental Changes of Translated Genes in RGC Axons, Related to Figure 4

(A) The upper heat map displays the enrichment of GO terms or KEGG pathways for axonally translated genes. Each row in the heat map corresponds to a single GO term. Genes are clustered either by stage-specific expression (“stages”) or hierarchical clustering (lower heat map) according to their developmental changes (“changes”).

(B) Ingenuity pathway analysis (IPA) to identify canonical pathways associated with the axonal translome. Each row represents a single pathway (blue, enriched). The right panel shows lists of pathways extracted from each cluster.

(C) A heatmap showing the log2 (read count) for adult samples with and without ribosome run-off. Each row in the heat map corresponds to a single gene.
Figure S5. Analysis of trans-Acting Elements that Regulate the Axonal Translatome, Related to Figure 5

(A) Ingenuity pathway analysis (IPA) to identify upstream regulators associated with the axonal translatome. The abundance of each mRNA between two consecutive stages was represented as the ratio (ratio > 1 indicates increase in translation). The coordinate change in the translation ratios was calculated as the activation z-score. A positive z-score indicates that the translational regulator is expected to be activated.

(B) Bar graph representing the fold change in levels of axonally translated genes. The mRNA levels in the axonal translatome were quantified by qRT-PCR (normalized by TRAPed cDNA for each stage).

(C) Density plots showing the distribution of changes in FPKM values for the axonal translatome during two consecutive developmental stages with p values (Komogorov-Smirnov test). The values are calculated as follows: log2 (stage A(FPKM)/stage B (FPKM)). The distributions of target genes in pathways, which are indicated by colored lines, are overlapped with non-target genes represented by gray lines.

(D) A scatter plot of the Principal Component Analysis (PCA) based on normalized read counts in the axonal and retinal translatome from four different stages. Data were plotted using the first two Principal Components (PCs), which explained up to 73.2% of the total variance.

(E) Relationship between transcript abundance of the genes in E17.5 transcriptome, which were detected in E17.5 axonal translatome, and probability of their translation at later stages (blue line, mRNA level in transcriptome; red line, moving averages of percentage of genes detected at any of three later stages over a window size of 100 genes; r, Pearson correlation coefficient). The right heatmap shows mRNA abundance in the translatome (left) and enriched pathways (right).
A  *Citα* isoforms (exon-skipping)

B  Axon-specific back splicing events

Figure S6. Analysis for Alternative Isoforms and *cis*-Acting Elements, Related to Figure 6

(A) Sequence reads (gray bars) mapped on the *Citα* gene. The mapped reads are visualized with Integrative Genomics Viewer (IGV).

(B) Sequence reads (gray bars) mapped on the *RhoBtb3* (upper panel). Sanger sequencing of RT-PCR fragment of the *Ankrd12* mRNA (lower panel).
Figure S7. Cis-Regulatory Elements for Axonal Translation, Related to Figure 7
(A) Lists of sequence motifs enriched in 5'UTRs, 3'UTRs and alternative exons of axon-enriched mRNAs / exons.
(B) An example of axon-enriched motifs. The scatterplot compares the normalized mRNA levels (log2(FPKM)) between the axonal (y axis) and the retinal (x axis) translatome at stage P0.5 for genes with (red dots) and without (black dots) the motif. The density plot shows the distribution of log2 (axon (FPKM) / retina (FPKM)).
(C) GO enrichment analysis for entire genome containing axon-specific sequence motifs (K: G or T; R: A or G; Y: C or T; M: A or C; R: A or G; and H A or C or T) and their relative efficiency in axonal mRNA translation measured by fluorescence recovery after photobleaching (FRAP) of motif-containing reporter constructs.

(legend continued on next page)
Several axon-specific motifs were able to promote mRNA translation in the growth cone relative to a control myr-d2EGFP construct without a UTR. Statistical significance of FRAP compared to the no-UTR control was tested across all time-points (0-10mins) using a two-way ANOVA (from the top bar, n = 16, 5, 5, 7, 8, 7, 3, 8, 8, 5, 8, and 6, respectively). For representative purposes, the mean fluorescence recovery at 10 min post-photobleaching is shown. Error bars represent SEM. **p < 0.01, and ***p < 0.001 compared to no-UTR control.
Supplemental Information

Dynamic Axonal Translation in
Developing and Mature Visual Circuits

Toshiaki Shigeoka, Hosung Jung, Jane Jung, Benita Turner-Bridger, Jiyeon Ohk, Julie Qiaojin Lin, Paul S. Amieux, and Christine E. Holt
SUPPLEMENTAL INFORMATION

The dynamic translatome of retinal ganglion cell axons during assembly and maintenance of the mouse visual system

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EXTENDED EXPERIMENTAL PROCEDURES

Animals

RiboTag and Pax6-alpha-Cre mice were kind gifts from Dr Paul Amieux (University of Washington) and Dr Peter Gruss (Max Planck Institute), respectively. A homozygote RiboTag female mouse was mated with a Pax6-alpha-Cre male, to produce Cre-positive and Cre-negative mice in a single litter. Cre-negative embryos or pups were used as negative controls for TRAP. Rosa26-StopLox-LacZ and Rosa26-StopLox-TauLacZ were kindly provided by Dr Jin-Woong Bok (Yonsei University) and Dr Soochul Park (Sookmyung Woman’s University), respectively. All procedures were conducted under license in accordance with UK Home office guidelines and under the Guidelines for the Care and Use of Laboratory Animals of Yonsei University College of Medicine.

Monitoring of Cre-recombinase activity in Pax6-alpha-Cre mice

As even slightly leaky expression of the HA-tagged ribosome in any SC-resident cells would lead to misidentification of axonally translating mRNAs, we needed to confirm that no resident cells in the SC express Cre. We took two independent approaches. In the first, we used two Cre-responsive reporter mice: one that labels the cell bodies of Cre-expressing cells and their progeny (Rosa26-StopLox-LacZ) and the other that labels the axons of these cells (Rosa26-StopLox-TauLacZ). In accordance with the previous reports (Marquardt et al., 2001), LacZ-positive cell bodies were only observed in the neural retina but not in the SC (Fig. S1). Using the StopLox-TauLacZ reporter mouse to visualize retinal axons, we could see the SC is richly innervated by the axons originating from their Cre-positive cell bodies in the retina (Fig. S1). In the second approach, we crossed this mouse with RiboTag (Fig. 1C), and asked whether we could detect any trace of the recombined RiboTag allele (HA-rpL22) in the SC. If the SC contains any resident cells that have expressed Cre but escaped our
histological analysis, the nuclear DNA extracted from this tissue must contain the
HA-\textit{rpl22} allele, which we can detect using PCR-based assays (Fig. 1D). We
detected no such signal in the SC dissected for TRAP (Figs. 1D and S1, red box).
Therefore, both histological and molecular biological assays confirm that the only
source of HA-tagged ribosomes in the SC is the RGC axons. Genotyping was
performed by polymerase chain reaction (PCR) using the following primer pairs: for
the Cre transgene, forward 5'- GCATTACCGGTCGATGCA ACGAGTG-3', and
reverse 5'- GAACGCTAGAGCCTGTTTTGCACGTTC-3'; for the RiboTag allele,
forward 5'- GGGAGGCTTGCTGGATATGACGAGTG -3', and reverse 5'- TTTCCAGACACAG-
GCTAAGTACAC-3'; for detection of the recombined RiboTag allele (HA-rpL22),
forward 5'- TTCTCTAGAAAGTATAGGAACG -3', and reverse 5'- ACATCGTGATGGG-
TATAGATCC-3'. Cre-negative embryos or pups were used as negative controls for
TRAP.

\textbf{Quantitative reverse transcription-polymerase chain reaction (qRT-PCR)}
cDNA was synthesized using an oligo-dT primer and SuperScript III reverse
transcriptase (Invitrogen). The qPCR was performed using QuantiTect SYBR Green
PCR Kit (Qiagen) on Light Cycler LC480 II (Roche). The levels were normalized by
the total RNA amounts. The following primer pairs were used: for Glud1, forward 5'-
GGGAGGTCATCGAAGGCTAC-3', and reverse 5'- AGCCAGTGCTTTTACTTCAT-
CC-3'; for Mapt, forward 5'- TTCTGTCTCCTCGCCTTCT-GTC-3', and reverse 5'-
CCTTCTTGGTCTTGAGGACG-3'; for Rps5, forward 5'- TCAAGCTCTTTGGGAAAT-
GG-3', and reverse 5'- GGGCAGGTACTTGGGATCT-3'; for Tsc2, forward 5'-
TAGGGCTCTGGTCTACCTT-3', and reverse 5'- GTGCT-TGTAATGGAGCTGA-3'; for Cfl1, forward 5'- TCTGTCTCCTCGGCGTCG-3', and reverse 5'-
GCCTTTCTGCTTCTTCCAC - 3', for Aldoa, forward 5'- TTAGTCTTTTGCCTAC-
CCCA-3', and reverse 5'- AGCTCTTTCTTGCTCGG-3'; for Atp5b, forward 5'-
CACAATGCAGAAAGGATCA-3', and reverse 5'- GGTCAGTCAGG-TCATCAGC-
3'; for Basp, forward 5'-ACAAAGACAAAGAGCCGAA-3', and reverse 5'-CTCTCCTTGACCTCGGTGG-3'; and for Cend1, forward 5'-CCTGAGCACTCCTCGGTATC-3', and reverse 5'-AGACCACAGTGGCTCAGGAC-3'.

**Histological analysis**

Mouse embryos were fixed by immersion in 4% paraformaldehyde in PBS. Adult mice were transcardially perfused with the same fixative, and entire eyes and brains were dissected out and post-fixed. For X-gal staining, brains and retinas were dissected out, washed in ice-cold PBS and fixed for 1 hour in 1% formaldehyde, 0.2% glutaraldehyde, 2mM MgCl₂, 5mM EGTA, 0.02% NP-40 in PBS (pH 7.5) at 4°C on a shaker. The dissected tissues were rinsed three times for 20 min each in PBS and stained with X-gal staining solution (1mg/ml X-gal diluted in DMF, 0.01% Sodium deoxycholate, 0.02% NP-40, 5 mM potassium ferricyanide, 2 mM MgCl₂ in PBS) between 3 to 48 hours at 37°C in the dark. After washing with PBS, the tissues were post-fixed in 4% paraformaldehyde in PBS for 30 min at 4°C. For immunohistochemistry, tissues were saturated in 30% sucrose, embedded in OCT, frozen on dry ice, sectioned at 12 µm by using a cryostat (Leica CM3050S), and then visualized using an anti-HA antibody (Abcam ab9110) and a secondary antibody conjugated to Alexa Fluor 488 (Life technologies). For immuno-gold electron microscopy, tissues were fixed in 4% paraformaldehyde in 0.1M HEPES (pH 7.4), and the HA-tagged ribosomes were visualized by the same anti-HA antibody and IgG conjugated with gold (10nm-15nm) at the Cambridge Advanced Imaging Centre (University of Cambridge, UK). Specific labeling was evident as ultrastructurally identifiable ribosomes were labeled by multiple gold particles (Fig. 1F, purple arrows). We did, however, occasionally observe scattered gold particles in the Cre-negative tissue (Fig. 1G, white arrow), but these were never clustered as seen in the Cre-positive group. Therefore, we considered only two or more gold particles in close
proximity (within 50 nm between particles) as specific labeling of HA-positive ribosomes.

**Axon-Translating Ribosome Affinity Purification (TRAP)**

Tissue samples were dissected and snap-frozen in liquid nitrogen and genotyped. Whole eyes were used as the cell body group and the superior colliculus, where retinal axons terminate, were used as the axon terminal group. Tissues from cre-positive and negative were pooled for TRAP (three eyes and superior colliculi for one group). Tissues were homogenized in lysis buffer (20mM HEPES-KOH, 5mM MgCl₂, 150mM KCl, 1mM DTT, SUPERase In, and Complete EDTA-free Protease Inhibitor Cocktail) in the presence of cycloheximide (to stop translational elongation and to lock translating ribosomes on the mRNA) and rapamycin (to prevent new translational initiation during immunoprecipitation), and post-mitochondrial fractions were collected. We optimized the TRAP protocol before performing axon-TRAP. First, we found the polyclonal HA antibody (9110, Abcam) is superior to the one used in the original RiboTag study (HA11, Covance) (Sanz et al., 2009). Although the two antibodies were similarly effective in precipitating HA-rpL22, the polyclonal antibody (Abcam ab9110) co-purified much more 80S ribosomes (Fig. 2A). We estimated that approximately 40% of HA-tagged translating ribosomes could be purified this way, as the amount of TRAPed 80S ribosomes (i.e. co-immunoprecipitated rpS3a, a 40S ribosomal protein) was approximately 10% of the input (i.e. rpS3a in total input) (Fig. 2A). This estimation is based on the findings that (1) roughly 50% of retinal cells express HA-tagged rpL22 (i.e. 50% cells do not express tagged ribosomes), and that (2) these cells express rpL22 from one wildtype allele and one recombined HA-rpL22 allele (i.e. 50% of rpL22 is labeled with HA in Cre-positive cells) ([total 80S] = [HA-80S] x (1/50%) x (1/50%) = [TRAPed HA-80S] x (1/10%). Therefore, [TRAPed HA-80S] = (4/10) x [HA-80S]) (Fig. 2A). We found that the indirect immunoprecipitation protocol, which allows the antibody to bind the antigen before the purification of the
antigen-antibody complexes, was more specific than the direct protocol, which utilizes the antibody pre-conjugated to Protein G-magnetic beads. We think that nonspecific binding of mRNAs to Protein G-magnetic beads was reduced in the indirect protocol, as the tissue lysate was pre-cleared with unconjugated Protein G-magnetic beads. Pre-cleared ribosome-mRNA complexes were immunoprecipitated by an anti-HA antibody and Dynabeads Protein G (Life Technologies 10004D). Total RNA was extracted from the ribosome-mRNA complexes using an RNeasy mini kit (Qiagen) followed by in-column DNase treatment to remove genomic DNA contamination. The RNA samples were examined for quantity and quality using the Agilent Bioanalyzer 2100 (Agilent Technologies).

**Amplification of cDNAs**

The amount of axon-TRAPed mRNA was minute and had to be amplified (Fig. S2A). We used a method developed by Tang and colleagues for single cell transcriptomics (Tang et al., 2009) with slight modification (Fig. S2A). First, TRAPed RNAs were treated with DNase I to eliminate genomic DNA contamination and then mRNAs were reverse-transcribed with oligo(dT) primer with a linker. After second strand synthesis with another linker, the double-strand cDNAs were amplified by ten rounds of PCR and visualized by agarose gel electrophoresis (Fig. 2B). The specificity of TRAP was evident because it was dependent on Cre. As expected, the amount of TRAPed mRNAs was lower in the SC (mRNAs from RGC axons) than in the eye (mRNAs from the cell body and proximal neurites), and therefore was further amplified by five additional rounds of PCR. Although axon-TRAP was clearly dependent on Cre and therefore specific, increasing PCR cycles led to an increased background (i.e. amplified cDNAs from the Cre-negative SC). We reasoned that the amplified DNAs in the Cre-negative SC would reflect the relative abundance of mRNAs in the SC. We sequenced these mRNAs and used the data as a negative control for bioinformatics analysis of axon-TRAPed mRNAs (Fig. 2B). Amplified
cDNAs were subjected to paired-end 100 or 90 bp sequencing on the Illumina HiSeq2000.

For the in vitro ribosome run-off experiment, eyes or superior colliculi were homogenized by the 400 µl lysis buffer without cycloheximide, and then 200 µl of rabbit reticulocyte lysate (Promega) and 8 µl of Harringtonine (100 ug /ml) and 4 µl of 4E1RCat (5 mM) were added to the 188 µl lysate, followed by 37 °C incubation for 30 min. To stop the in vitro translation elongation, 800 µl of ice-cold lysis buffer with cycloheximide was added, followed by immunoprecipitation with the anti-HA antibody. The purified RNAs were subjected to RNA sequencing on the Illumina NextSeq 500. To reflect the difference in amount of amplified cDNAs between the samples with and without run-off translation on the sequencing depth, we first measured the amount of cDNA by both Tapestation DNA and by Qubit. We then adjusted the ratio between the amounts of input libraries for multiplex sequencing using the ratio of cDNA amount between the two samples. All RNA-seq data are deposited in Gene Expression Omnibus (GEO) datasets under accession number GSE79352.

**Mapping of sequence reads and normalization of read counts**

The sequence reads were mapped to the mouse genome (mm10) using TopHat 2 version 2.0.12 (Kim et al., 2013) with default settings, except for the "--read- realign-edit-dist 0" option. This option was chosen to reduce mapping to pseudogenes. Although we detected a significant number of sequencing reads of the primer dimers in several samples, which are probably caused by the matching problem between our primers and Illumina primers, because there is no biological reason that these dimers bias the relative abundance of sequence reads of endogenous mRNAs, we analyzed all sequence reads that can be mapped to the mouse genome sequence by TopHat 2. Transcript assembly and estimation of FPKM (Fragments Per Kilobase of
transcript per Million fragments sequenced) values were performed using Cufflinks version 2.2.1 (Trapnell et al., 2010). Read counts for each gene were determined using HTSeq version 0.6.1p1 (http://www-huber.embl.de/users/anders/HTSeq/). For the analysis of run-off samples, we used the read counts for all DEGs (adult) detected either in the run-off positive or negative sample.

**Differential gene expression analysis and Gene Ontology based enrichment analysis**

For the identification of translated mRNAs in RGC axons, we applied the differential gene expression analysis on read count data between two biological replicates of Cre-positive and Cre-negative samples using NOISeq (Tarazona et al., 2011) in default conditions with probability threshold 0.7. Although this approach has a risk of filtering out actively translating mRNAs if the same mRNAs exist in high abundance in SC-resident cells and are proportionately represented in the negative control, we thought that it would be appropriate to take a conservative approach when analyzing mRNAs identified by highly sensitive RNA-seq from samples with potentially low signal-to-noise ratio. DEG analysis identified a subset of Cre-dependent mRNAs (Fig. 2D, right panel, red dots) (Table S1). In some cases, genes that passed our filtering criteria and were identified as DEGs in one stage failed the same test in other stages (Fig. 2D, orange), because their abundance in the negative control varied depending on the stage. Most of these mRNAs, however, were found in the axon-TRAPed mRNAs at all stages tested, suggesting that these genes became false negatives in other stages. Therefore, when we analyzed the developmental change in translation levels, we used the FPKM (fragments per kilobase of transcript per million mapped reads) values of all four stages for genes that have passed the DEG test in at least one stage (Union of DEGs) (Fig. 2D, orange and peach).
To perform a GO-based analysis for neuronal functions (Figs. 4A and S4A), we selected 455 neuron-related GO terms (Table S2) using the following criteria: all offspring GO terms (249 terms) of 'Neuron development (GO:0048666)', which contain GO terms related to neuron/axon development, and all offspring GO terms (206 terms) of "Synaptic transmission (GO:0007268)", which contain GO terms related to neurotransmission. The offspring GO terms were identified using the GOBPOFFSPRING function of GO.db, an R package (Table S2). The enrichment analysis for gene ontology was carried out with DAVID and topGO version 2.18 (Alexa et al., 2006) on R version 3.1.2. The result for all analyzed GO terms (biological process) is represented in Table S3. For Kyoto Encyclopedia of Genes and Genomes (KEGG) analysis, we used all 2576 axonally translated genes and calculated p values and fold enrichments using DAVID (Fig. 4A, lower panel). The numbers of GO terms associated with retinal and axonal translatome were analysed using ClueGO (Bindea et al., 2009). Statistically significant enrichment is annotated by a solid box outline.

**Ingenuity Pathway Analysis**

Ingenuity Pathway Analysis (Qiagen) for canonical pathways (Fig. 5A) was performed according to the manufacturer’s instructions. For quantitative analysis of stage-dependent axonal translation of each mRNA, we calculated the ratio in read counts of each mRNA from one stage to the next. Then we performed Ingenuity Pathway Analysis (IPA) for upstream regulators, which mainly utilizes published results of gene knockdown or knockout studies. For example, if the protein product of a gene decreases when a specific translational regulator is knocked out, the gene would be described as “positively regulated” by the translational regulator (and vice versa) (Fig. 5B).

**Analysis of target mRNAs in the mTOR pathway and RNA binding proteins**
We analyzed the known binding targets of RBPs using the results of previous studies on FMRP, TDP-43, FUS and APC. In addition, we carried out the same analysis on the targets of mTORC1 to confirm the result of IPA analysis. If any of these molecules regulate stage-specific translation in the axon, their target mRNAs would show coordinate changes in translation. We used the gene sets that were identified in previous studies (Colombrita et al., 2012; Darnell et al., 2011; Preitner et al., 2014; Thoreen et al., 2012). We analyzed mTOR targets that are described in the previous report (Thoreen et al., 2012) with a threshold of log2 (Torin1/Vehicle) < -1, FMRP targets (Darnell et al., 2011) with a threshold of rank (based upon chi-square score) < 100 and all of the APC targets (Preitner et al., 2014), TDP-43 targets and FUS targets identified (Colombrita et al., 2012). In the FPKM ratio analysis for consecutive developmental stages, we used all genes that were detected in RNA-seq in order to avoid any bias caused by differential gene expression analysis. Principle component analysis (PCA) was performed on normalized read counts of all samples to compare gene expression in 12 different conditions by using the “prcomp” function in the R Software package (version 2.13.0). Data were plotted using the first two PCs which explained up to 73.2% of the total variance: 66.1% explained by PC1; and 7.1% by PC2.

**Analysis of mRNA isoforms**

Alternative isoforms were analyzed on mapped reads from the P0.5 sample using MISO. Before this analysis, the mean and the standard deviation of the insert length and the total number of mapped read pairs were computed using the “pe_utils” utilities (Retina: mean=177.8, sdev=9.7, dispersion=0.7, num_pairs=3262858, Axon: mean=178.3, sdev=13.5, dispersion=1.0, num_pairs=2535309, Cre_negative axon: mean=179.6, sdev=9.2, dispersion=0.7, num_pairs=3235484). We used Mouse genome (mm10) alternative events version 1.0 (Wang et al., 2008) to perform “exon-centric” analyses. We counted and analyzed only the events in which both splicing
variants are detected both in axon and in cell body (the events of “0 < Ψ <1” both in axon and in cell body). For the discovery of novel splice variants, we performed reference-independent transcript reconstruction using Cufflinks version 2.2.1 (Trapnell et al., 2010) and compared the sequences of reconstructed transcripts with the UCSC Genes transcript annotations by using the Cuffcompare program from the Cufflinks package. For the identification of circular RNAs, we first extracted the fusion transcripts from the unmapped sequence reads by using TopHat-fusion (Kim and Salzberg, 2011) (TopHat 2.0.12, parameters:--fusion-search --keep-fasta-order --bowtie1 --no-coverage-search), and then identified junction reads from back spliced exons using CIRCexplorer-1.1.1(Zhang et al., 2014). Mapped sequence reads are visualized using the Integrative Genomics Viewer (IGV) version 2.3.46 (Robinson et al., 2011).

**Identification of regulatory motifs in RNA**

*De novo* motif analysis was performed using HOMER version 3.0 (Heinz et al., 2010) with custom FASTA files. For the finding of motifs in UTRs, we used the UTR sequences of genes whose levels were higher in the axon than in the retina (axon / retina > 100, Fig. 3A) in P0.5 translatomes. For finding of motifs in axon-enriched alternative exons, detection of differential exon usage was performed using DEXseq version 1.10.8 (Anders et al., 2012) on R version 3.1.0, and exons were selected with a Benjamini–Hochberg adjusted p-value cut-off of < 0.1 (Benjamini and Hochberg, 1995). To validate these motifs, we extracted the genes that contain these motifs allowing 0-1 mismatch and then compared the translation levels between axon and retina. Sequences of UTRs were retrieved from Ensemble BioMart (Kinsella et al., 2011). Motif containing genes were identified using the Biostrings package on R version 3.1.2.

**Immunofluorescence**
Mouse RGC axons were cultured as previously described (Zivraj et al., 2010). The cultures were fixed in 2% (vol/vol) paraformaldehyde/7.5% (wt/vol) sucrose, permeabilized with 0.1% Saponin (Sigma) and blocked in 5% goat serum, then labeled with primary antibodies overnight and Alexa Fluor 488 secondary antibodies (1:1000, Life Technologies), and mounted in FluorSave (Calbiochem). Randomly selected isolated axons were imaged with a Plan Apo 60X oil objective on a Nikon Eclipse TE2000-U inverted fluorescent microscope with a Hamamatsu ORCA-ER CCD camera. Exposure time was kept constant and below greyscale pixel saturation. For quantitation of fluorescence intensity, the outline of the axon segments with similar lengths was traced on the phase image using Volocity software (PerkinElmer), and then superimposed on the fluorescent image. The software calculated the fluorescent intensity within the axon segment, giving a measurement of pixel intensity per unit area. The axon outline was then placed in an adjacent area clear of cellular material to record the background fluorescent intensity. This reading was subtracted from the axon reading, yielding the background-corrected intensity. Each reading was normalized to the mean fluorescent intensity of E17.5 axons in the same group. The fluorescent intensities of between 50 to 100 axon segments per sample group were collected. The following antibodies were used: anti-mTOR (phospho S2448) antibody (Abcam 109268), anti-FMRP antibody (Abcam 17722), and anti-RPS6 (phospho S235 + S236) antibody (Abcam 12864).

**Fluorescence recovery after photobleaching (FRAP)**

Motif sequences were cloned into a plasmid containing the coding sequence of membrane-targeted, destabilized EGFP (myr-d2EGFP) (Aakalu et al., 2001) between NotI-Xhoi sites (for 3'-UTR and alternative exon) or Nhel-BamHI sites (for 5'-UTR). Motif sequences are in Table S3. The reporter plasmids were expressed in the retina by targeted electroporation into *Xenopus laevis* embryos, and fluorescence was imaged in cultured retinal ganglion cell axons as previously described (Leung et
al., 2006), except that 1% fetal bovine serum was added to the medium. In experiments to confirm translation-dependency of FRAP, the translation inhibitor anisomycin (40 µM) was added 30 min prior to imaging. FRAP experiments were performed on an Olympus IX81 inverted microscope equipped with a PerkinElmer Spinning Disk UltraVIEW VoX and a 60x (NA, 1.30) Olympus silicone oil immersion objective. Images were acquired with an ORCA-Flash4.0 V2 CMOS camera (Hamamatsu) using Volocity software (PerkinElmer).

Photobleaching was performed using an UltraVIEW PhotoKinesis device (PerkinElmer). Regions of interest were manually defined so that growth cones and >20µm of the axon shaft were bleached (thus reducing likelihood of fluorescence recovery resulting from myr-d2EGFP diffusion from unbleached areas of the axon shaft). Photobleaching was performed at 50-85% laser power (488 nm laser line) with 20–30 bleach cycles. Time-lapse images were captured at 1 min intervals using a 488 nm laser line at 25% laser power for myr-d2EGFP and a 561 nm laser line (31 % laser power) for visualization of axons using a membrane-targeted mCherry-CAAX reporter, in addition to phase contrast. Exposure time for the 488 channel was adjusted to avoid pixel saturation and was typically between 50-200ms.

Quantification of fluorescence intensity was performed using Volocity software (PerkinElmer). At each time point, the outline of the growth cone (ROI) was traced using the mCherry-CAAX reporter (561 channel) and phase contrast images. Mean gray values from the 488 channel were subsequently calculated as mean pixel intensity per unit area within the ROI. Subtracting the mean background pixel intensity per unit area from an equivalently sized ROI immediately adjacent to the growth cone normalised intensity values in the growth cone. Unhealthy axons exhibiting signs of photo-toxicity after FRAP (characterised by blebbing, growth cone collapse and/or retraction) were excluded from analysis. In addition, we only quantified growth cones of axons extending more than 100 µm from the eye explant.
to reduce effects of somal diffusion. Relative fluorescent recovery (R) at each time point was calculated by the formula: \( R_x = (I_x - I_{\text{post}}) / (I_{\text{pre}} - I_{\text{post}}) \). Where, \( I_x = \) normalised fluorescent intensity of the growth cone ROI at time point ‘x’, \( I_{\text{pre}} = \) normalised fluorescent intensity before photobleaching and \( I_{\text{post}} = \) normalised fluorescent intensity immediately after photobleaching (t=0 mins).

Data were analysed using PRISM software (Graphpad). Significance was tested using a two-way ANOVA.

REFERENCES


