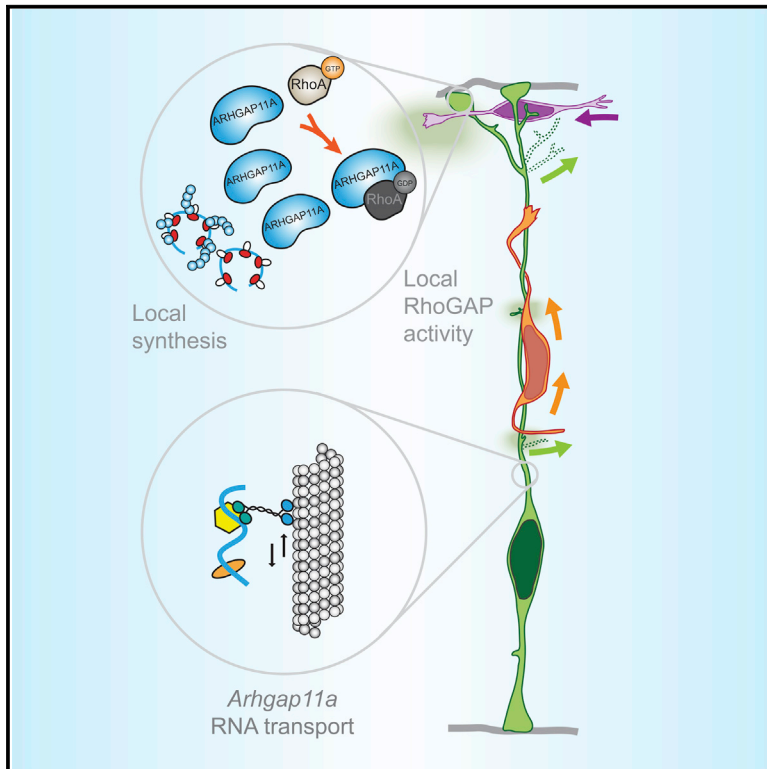


Subcellular mRNA localization and local translation of *Arhgap11a* in radial glial progenitors regulates cortical development

Graphical abstract



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In brief

Pilaz et al. demonstrate that *Arhgap11a* mRNA undergoes active transport and local translation in radial glial basal endfeet, which is essential for proper endfeet morphology. Further, *Arhgap11a* non-cell autonomously impacts positioning of interneurons and excitatory neurons. This demonstrates the functions of local translation in radial glial progenitors.

Highlights

- *Arhgap11a* subcellularly localizes in radial glia of mouse and human cerebral cortex
- *Arhgap11a* 5' UTR directs active transport and local translation in radial glia
- Localized mRNA and RhoA-GAP activity controls radial glial endfoot morphology
- *Arhgap11a* non-cell autonomously acts in radial glia to mediate interneuron position



Article

Subcellular mRNA localization and local translation of *Arhgap11a* in radial glial progenitors regulates cortical development

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SUMMARY

mRNA localization and local translation enable exquisite spatial and temporal control of gene expression, particularly in polarized, elongated cells. These features are especially prominent in radial glial cells (RGCs), which are neural and glial precursors of the developing cerebral cortex and scaffolds for migrating neurons. Yet the mechanisms by which subcellular RGC compartments accomplish their diverse functions are poorly understood. Here, we demonstrate that mRNA localization and local translation of the RhoGAP ARHGAP11A in the basal endfeet of RGCs control their morphology and mediate neuronal positioning. *Arhgap11a* transcript and protein exhibit conserved localization to RGC basal structures in mice and humans, conferred by the 5' UTR. Proper RGC morphology relies upon active *Arhgap11a* mRNA transport and localization to the basal endfeet, where ARHGAP11A is locally synthesized. This translation is essential for positioning interneurons at the basement membrane. Thus, local translation spatially and acutely activates Rho signaling in RGCs to compartmentalize neural progenitor functions.

INTRODUCTION

In eukaryotes, subcellular RNA localization and local translation allow cells to temporally and spatially control functions that rely on dynamic and complex proteomes. In highly polarized cells, such as neurons and migrating fibroblasts, mRNA localization plays a pivotal role in local cytoskeletal regulation and hence local morphology.^{1,2} Most localization studies use cultured cells or mechanical axotomy, limiting an understanding of events *in vivo*. Notably, the developing and adult brains contain some of the most highly polarized and elongated cell types found in animals. These features are especially prominent in radial glial cells

(RGCs), which control cortical development by acting as neural stem cells to generate neurons and then astrocytes and by scaffolding radial neuron migration.^{3–8}

RGCs are bipolar, with basal processes emanating from cell bodies in the ventricular zone (VZ) and radially traversing the cortex to form basal endfeet at the pia and apical endfeet at the ventricle.^{9,10} The basal process can be extremely long, reaching several hundred microns in the mouse and centimeters in humans. The apical and basal endfeet encounter unique niches, with the latter including interneurons, Cajal-Retzius neurons, and excitatory neurons.¹¹ Basal endfeet are tightly connected to the basal lamina, forming a barrier between the brain and



the overlying meninges, composed of fibroblasts and blood vessels.^{12–14} Along the basal process and at endfeet, dynamic filopodia-like protrusions extend and retract, which is postulated to influence signaling and neuronal migration.¹⁵ Further, as development proceeds, basal endfeet become more complex in number¹⁶; basal structures are also notably complex in humans.^{17–19} Disruptions to RGC morphology can have disastrous consequences on the architecture of the mature cortex, causing cobblestone malformation and lissencephaly.^{13,20} Thus, RGC morphology and subcellular compartmentalization are central to cortical development. Yet, we know surprisingly little about the cellular and molecular mechanisms mediating the proper morphology, dynamics, and function of distal RGC basal structures.

RGC endfeet are major sites of mRNA localization and are enriched for transcripts encoding cytoskeletal and signaling regulators, including GTPase regulators.^{21,22} Notably, the ubiquitous Rho GTPase is essential for cortical development,²³ but whether and how modulation of localized Rho activity controls RGCs is unknown. *Arhgap11a* encodes a RhoA-specific GAP, which promotes GTP hydrolysis and therefore inactivates the small GTPase RhoA.²⁴ Hence, the loss of *Arhgap11a* is predicted to increase RhoA signaling. *Arhgap11a* has essential roles in modulating the cytoskeleton, including mediating cytokinesis,²⁵ cell invasion,^{26,27} and neurite outgrowth.²⁸ In this light, local synthesis of a Rho regulator, such as ARHGAP11A, could help dictate the subcellular morphology and function of RGCs. However, the requirement of local translation of any transcript in RGCs has never been examined.

In this study, we tested whether mRNA localization and local translation of the RhoGAP ARHGAP11A in endfeet mediate RGC morphology and cortical development. We show that both *Arhgap11a* mRNA and protein subcellularly localize to RGC basal endfeet and basal processes during cortical development in mice and humans. Consistent with a role in these structures, *Arhgap11a* depletion from RGCs disrupts basal processes and endfeet morphology. This causes a non-cell autonomous impairment of excitatory neuron migration and inhibitory neuron positioning. We show the *Arhgap11a* 5' UTR is critical for its active transport in the basal process and local translation in basal endfeet. Importantly, impaired RGC morphology is rescued only when *Arhgap11a* localizes to endfeet and when RhoGAP activity is intact. We further find that rescuing morphology restores the proper position of interneurons along the basement membrane. Altogether, our study establishes critical, new *in vivo* requirements for subcellular mRNA localization in neural stem cells of the developing brain.

RESULTS

***Arhgap11a* mRNA and protein subcellularly localize to RGC basal endfeet midway through cortical development**

In this study, we asked whether the morphology and functions of RGCs are mediated by subcellular targeting of specific mRNAs and their local translation (Figure 1A). From our prior study, we noted that RGC basal endfeet are significantly enriched for mRNAs encoding cytoskeletal and GTPase signaling regula-

tors.²¹ We therefore sought to investigate the functions of transcripts associated with these cellular processes. *Arhgap11a* stood out as an outstanding candidate, given its established role in cytoskeletal regulation, neurite outgrowth, and GTPase signaling.^{24,25,27,28}

To examine the role of *Arhgap11a* in RGCs, we assessed its expression during mouse cortical development. Cortical neurogenesis in mice occurs between E11 and E18.5 and in humans, at gestational week (GW) 7–24.^{29–31} We used E14.5 *Dcx*-DsRed transgenic mice³² together with FACS and quantitative PCR (qPCR) to measure *Arhgap11a* mRNA expression in both progenitors (DsRed-negative) and neurons (DsRed-positive) (Figures 1B and S1A). Notably, *Arhgap11a* was expressed in progenitors but absent from newborn excitatory neurons. Likewise, in single-cell RNA sequencing datasets of the developing mouse and human cortex, *Arhgap11a* was highly enriched in RGCs but absent from post-mitotic excitatory and inhibitory neurons (Figures S1B–S1D).^{33–35} These data demonstrate that *Arhgap11a* expression in the developing cerebral cortex is largely specific to progenitors, including RGCs.

Next, we used immunohistochemistry to evaluate the expression pattern of the ARHGAP11A protein in the developing mouse cortex. At E11.5, ARHGAP11A localized within the germinal zones (Figures 1C–1E). However, strikingly, beginning at E13.5, the ARHGAP11A protein became enriched near the pial surface, at presumptive RGC endfeet, and along basal processes (Figures 1C, 1D, and 1F). This pial localization of ARHGAP11A was especially visible at later stages (Figures 1C, 1D, and 1G). We verified ARHGAP11A expression and localization in RGCs by co-staining with the RGC intermediate filament marker NESTIN at E15.5 (Figure 1H). In contrast, ARHGAP11A did not co-localize with Reelin-positive Cajal-Retzius neurons at the pia and was not expressed in interneurons (Figures S1C–S1F). This timing of ARHGAP11A localization to RGC basal processes and endfeet coincides with the onset of increased branching of endfeet,¹⁶ suggesting that this RhoGAP could locally influence RGC basal morphology.

We next examined *Arhgap11a* mRNA localization over the course of corticogenesis, to determine if its spatial and temporal pattern matched that of the protein. Toward this end, we used traditional as well as single-molecule fluorescent *in situ* hybridization (smFISH). At E14.5, *Arhgap11a* mRNA was significantly enriched at the pial surface compared with the VZ (Figures 1I, S1G, and S1H). Reinforcing the specific expression of *Arhgap11a* in RGCs, *Arhgap11a* co-localized with EGFP-labeled RGCs (introduced 1 day earlier via *in utero* electroporation [IUE]) but was notably absent from Tuj1-positive neurons (Figures S1G and S1H). At E15.5 and E16.5, *Arhgap11a* localization at the pia was especially prominent. smFISH confirmed *Arhgap11a* mRNA expression in EGFP-labeled RGC basal endfeet (Figures 1K and 1L). *Arhgap11a* RNA and protein co-localized at the pia, further evidencing this expression pattern (Figure 1M).

Given this striking localization pattern of *Arhgap11a* to RGC endfeet and the conserved expression within RGCs of mice and humans (Figures S1B–S1D), we next tested whether *Arhgap11a* mRNA is also present in the basal endfeet region of the human developing cortex. Thus, we employed *in situ* hybridization on post-conceptual week 11 tissue sections.

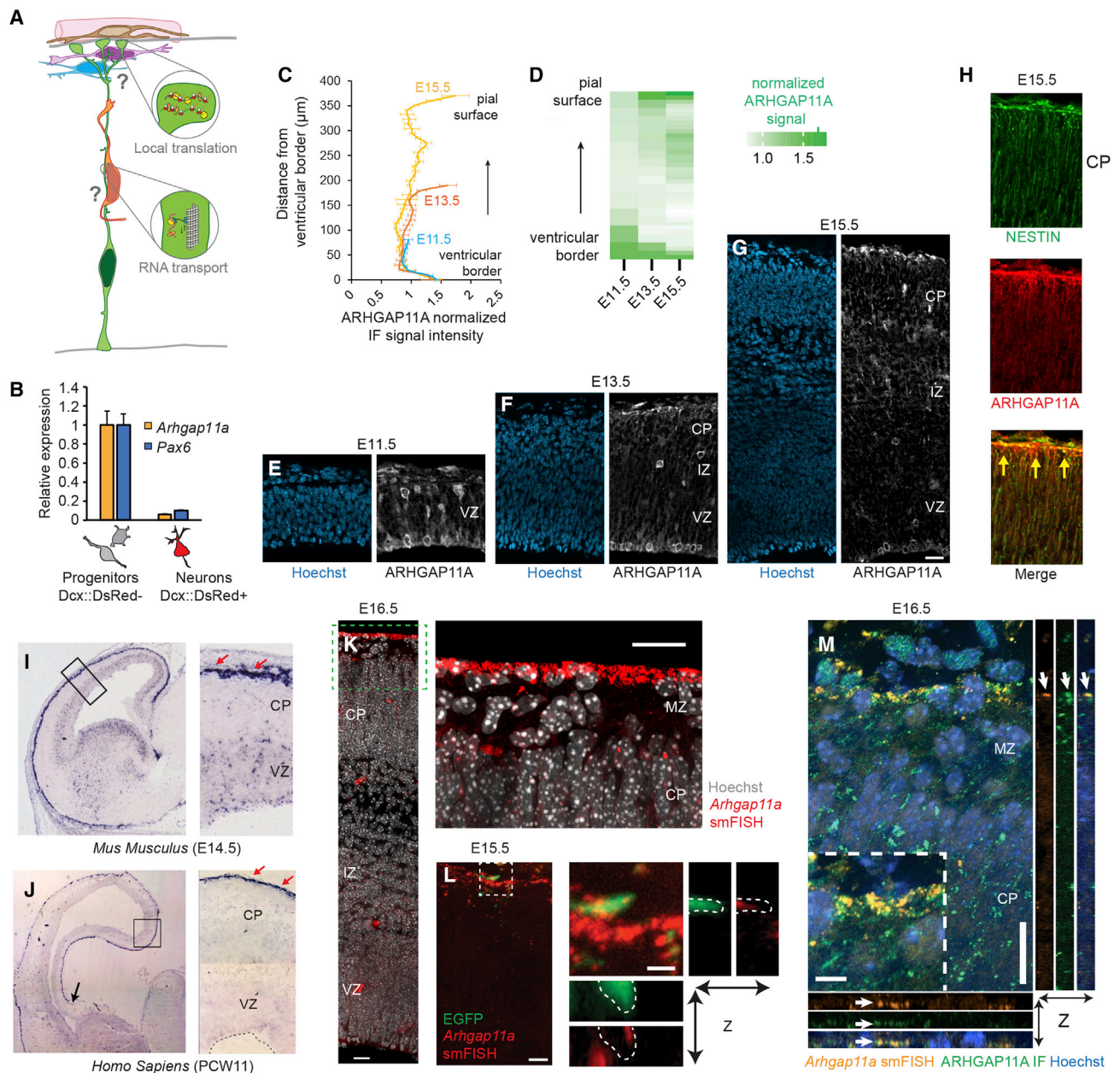


Figure 1. Subcellular localization of *Arhgap11a* mRNA and protein to RGC basal processes and endfeet during cortical development
 (A) Cartoon of a radial glial progenitor (RGC, green) with mRNA transport along the basal process and local translation in endfeet. Question marks reflect goal of the present study: what is the role of mRNA subcellular localization and translation in RGCs and for positioning of excitatory neurons (orange), migratory interneurons (purple), and Cajal-Retzius neurons (blue)?
 (B) qPCR analyses of *Arhgap11a* mRNA levels in E14.5 sorted embryonic cortical cells (n = 4 brains, 3 technical replicates).
 (C and D) Quantification of ARHGAP11A immunofluorescence in E11.5, E13.5, and E15.5 cortices.
 (E–G) Immunofluorescence of ARHGAP11A (gray) and Hoechst (blue) in E11.5 (E), E13.5 (F), and E15.5 (G) cortices.
 (H) Immunofluorescence of ARHGAP11A (red) at E15.5, showing expression in NESTIN positive RGCs (green) with overlap (yellow signal) in basal process and endfeet at the pial surface (yellow arrows).
 (I and J) *In situ* hybridization of *Arhgap11a* mRNA (purple signal), showing strong enrichment at the pia where RGC basal endfeet reside (red arrows) in E14.5 mouse brains (I) and in GW11 human fetal brains (J).
 (K and L) smFISH *in situ* hybridization depicting *Arhgap11a* mRNA (red) at the pia at E16.5 (K) and in EGFP+ RGC basal E16.5 endfeet (brains electroporated 1 day earlier) (L). Right panels, magnified areas highlighted in left panels (K and L) and maximum intensity projections of a z stack (L).
 (M) smFISH and immunofluorescence targeting *Arhgap11a* mRNA (red) and protein (green), respectively, highlights colocalization (arrows) in RGC basal endfeet. VZ, ventricular zone; CP, cortical plate; IZ, intermediate zone; smFISH, single-molecule fluorescent *in situ* hybridization. Scale bars: (E–G) 20 μ m; (K) 20 μ m; (L) left panel: 5 μ m, right panel: 1 μ m; (M) 20 μ m. Graph and bar plot: mean \pm SEM

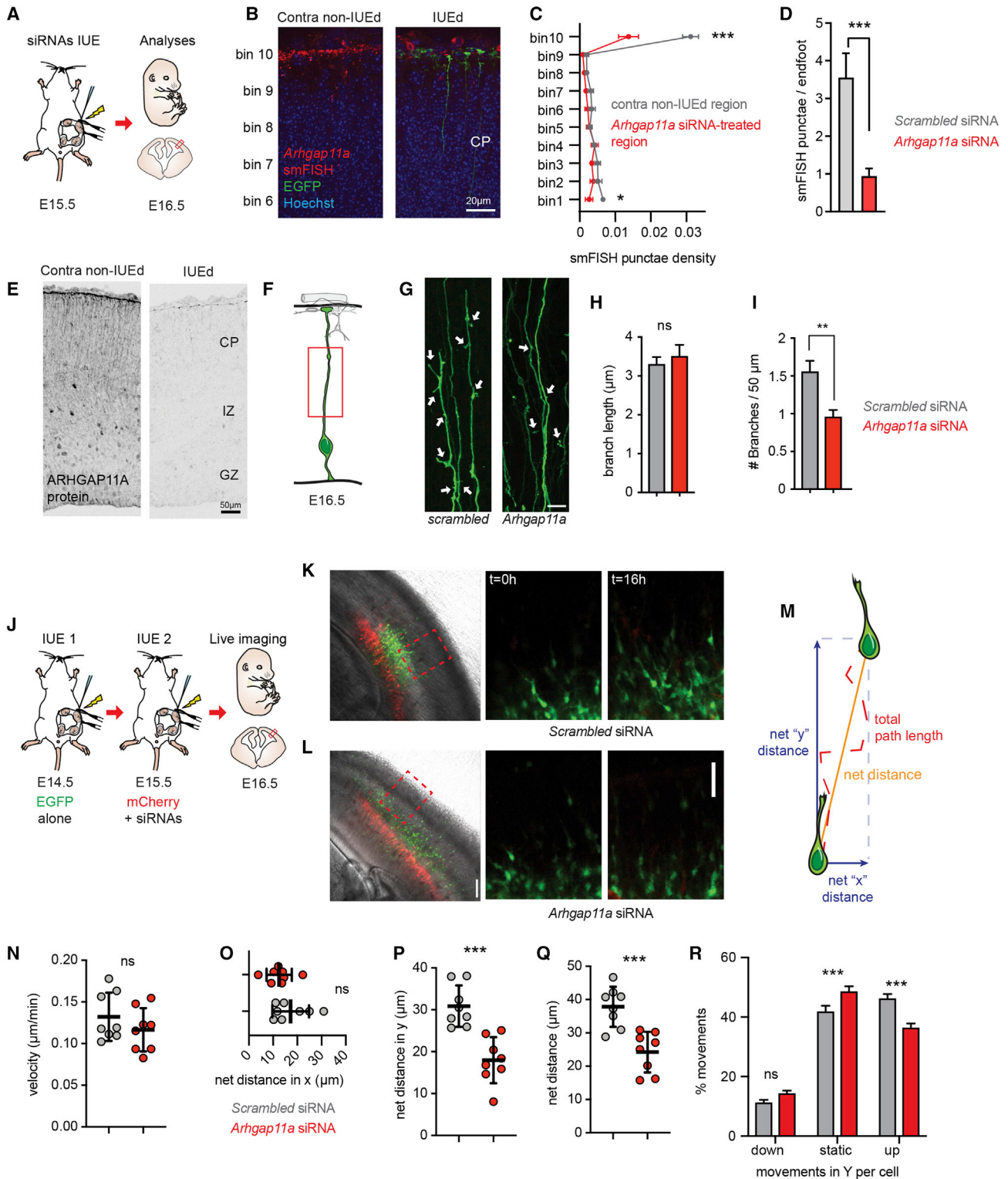


Figure 2. *Arhgap11a* controls RGC basal process morphology and non-cell autonomously controls radial migration of excitatory neurons (A) Schematic overview of the experiments in (B)–(I). (B–E) *Arhgap11a* mRNA is depleted from endfeet in the *Arhgap11a* siRNA electroporated region (IUE, green), evidenced by smFISH (red) (B) and immunofluorescence (E). (C) Binned quantification of *Arhgap11a* smFISH punctae in electroporated and contralateral non-electroporated regions. Bin 1 is apical lining the ventricle, and Bin 10 is adjacent to the meninges. (D) Quantification of *Arhgap11a* smFISH punctae in electroporated RGC endfeet.

(legend continued on next page)

ARHGAP11A mRNA exhibited conserved, robust enrichment at the pia in human fetal cortices (Figure 1J), in line with a previous report.³⁶ Altogether, these data demonstrate that *Arhgap11a* RNA and protein exhibit concordant subcellular localization in a developmentally controlled fashion to RGC basal processes and endfeet.

Arhgap11a is critical for RGC basal process morphology and non-cell autonomous control of radial neuronal migration

ARHGAP11A localization to RGC basal processes and endfeet coincides developmentally with the increasing morphological complexity of these structures.¹⁶ Further, excitatory neurons rely on the integrity of RGC basal processes to migrate to the cortical plate (CP).^{6,37,38} Given that ARHGAP11A has established functions in modulating RhoA signaling and cytoskeletal morphology,^{24,25,28} we hypothesized that *Arhgap11a* could regulate basal process morphology and thus influence excitatory neuron migration. To evaluate these possibilities, siRNAs targeting the 3' UTR of *Arhgap11a* mRNA were introduced by IUE together with a membrane-localized EGFP reporter driven by an RGC-specific promoter (pGLAST-EGFP-CAAX, Figure 2A). We manipulated *Arhgap11a* expression at E15.5, reasoning that alterations to basal process complexity should be evident at this stage.¹⁶ Using siRNAs, we effectively depleted *Arhgap11a* in RGCs both at the mRNA and protein levels, in RGC cell bodies and basal endfeet (Figures 2B and 2C, bins 1 and 10, respectively). Quantification of *Arhgap11a* RNA punctae per endfoot further validated this finding (Figure 2D). Furthermore, we observed little FISH signal in the intermediate zone (IZ) or CP within the control, thus reinforcing the specific expression of *Arhgap11a* in RGCs but not neurons. Finally, immunofluorescence showed a drastic reduction in ARHGAP11A protein levels in the electroporated area (Figure 2E).

We next examined the morphology of the *Arhgap11a*-depleted RGC basal process. For this, we generated 3D reconstructions of either scrambled (control) or siRNA-treated EGFP+ basal processes to quantify the density and length of extensions in the basal process (Figures 2F–2I). There was no impact of *Arhgap11a* depletion upon the length of small extensions (average extensions were 3.3 and 3.5 μm for control and *Arhgap11a* conditions, respectively) (Figure 2H). However, we

noted a significant decrease in the density of cellular extensions in brains transfected with *Arhgap11a* siRNAs compared with the control (Figure 2I). These data demonstrate that *Arhgap11a* regulates branching along the RGC basal process.

The integrity of the RGC basal process is paramount for proper radial migration of excitatory neurons from the VZ to the CP.^{6,37} Indeed, ectopic basal process branching can alter the patterns of migrating neurons, including speed, directionality, and pausing.³⁸ Therefore, we assessed whether aberrant RGC morphology induced by *Arhgap11a* loss impacts neuron migration in a non-cell-autonomous manner. To do this, we used a paradigm relying on sequential IUE on consecutive days (Figure 2J). E14.5 brains were IUE'd with an EGFP plasmid to label newborn cells that would become migratory neurons 1 day later (E15.5). At E15.5, we then performed an IUE in the identical location using scrambled or *Arhgap11a* siRNAs and membrane-bound mCherry to manipulate and label the RGC scaffold upon which EGFP+ neurons migrate. Importantly, since *Arhgap11a* is not expressed in migrating excitatory neurons (Figures 1 and S1), this allowed us to quantify the migration of WT neurons along an *Arhgap11a*-deficient RGC scaffold. At E16.5, we performed overnight (16 h) live imaging of EGFP+ migrating neurons along mCherry+ RGCs in organotypic brain slices, focusing on migration within the sub-ventricular zone (SVZ) and IZ (Figures 2K and 2L; Videos S1 and S2).

From live imaging, several parameters of neuronal migration were quantified: average speed, maximum speed, total distance, net distance, net distance in the y and x axes (in the radial or tangential dimensions, respectively), and fraction of time spent mobile, including toward the CP or VZ (Figures 2M–2R and S2A–S2G). Loss of *Arhgap11a* in RGCs had no impact upon average or maximum velocity, total path length (distance), or net distance of neurons traveled in the x axis (Figures 2N, 2O, S2C, and S2D). However, there was a significant 35% decrease in the net migrated distance and a particularly striking 42% decrease in the net distance on the y axis (Figures 2P and 2Q). Additionally, radially migrating neurons were more immobile (static) and showed a decreased propensity to move toward their final destination in the CP (Figure 2R). Thus, *Arhgap11a* loss does not impair the speed or ability of neurons to migrate per se but significantly reduces the efficiency and trajectory of radial migration. This was further corroborated by analysis of the

(F) Cartoon of regions analyzed in RGC basal processes.

(G) EGFP electroporated RGCs depicting reduced branches (arrows) along the basal process following *Arhgap11a* knockdown.

(H) Quantification of the length of branches along the RGC basal process (scrambled: n = 101 branches, 3 brains, 3 independent experiments; *Arhgap11a*: n = 72 branches, 3 brains, 3 independent experiments, unpaired t test with Welch's correction).

(I) Quantification of the density of branches along the RGC basal process (scrambled: n = 112 cells, 6 brains, 5 independent experiments, *Arhgap11a*: n = 99 cells, 5 brains, 4 independent experiments, unpaired t test with Welch's correction).

(J) Schematic overview of the experiments in (J)–(R) aimed at testing the impact of *Arhgap11a* depletion in RGCs on neuronal migration. Sequential IUEs were performed to label neurons (EGFP, green) and RGCs (red) at E16.5 when analysis is performed.

(K and L) Representative images showing electroporated regions (left) and position of migrating of neurons (green) at the beginning (t = 0 h, middle) and end of the live imaging experiment (t = 16 h, right).

(M) Neuronal migration parameters analyzed.

(N–R) Quantification of velocity of neuronal migration (N), net distance in X trajectory (O), net distance traveled in Y trajectory (P) and compiled distance (Q). (R) *Arhgap11a* knockdown in RGCs non-cell autonomously causes neurons to undergo more static movements and fewer movements toward the cortical plate (up) (scrambled and *Arhgap11a*: n = 8 brains, 2 independent experiments, unpaired t tests).

siRNAs, small interfering RNAs; IUE: *in utero* electroporation; CP, cortical plate; IZ, intermediate zone. *p value < 0.05. **p value < 0.01. ***p value < 0.001. Scale bars: (B) 20 μm , (E) 50 μm , (G) 10 μm , (K and L) right panels, 50 μm , left panels: 100 μm . Graphs and bar plots, means \pm SEM.

orientation of neurons at the beginning of the live imaging experiments, when EGFP+ neurons were mostly located in the SVZ and lower IZ. *Arhgap11a* loss increased the tangential orientation of those neurons (Figures S2E–S2G). Altogether, these results show that *Arhgap11a* expression in RGCs non-cell autonomously controls radial migration of pyramidal neurons along basal processes; this broadly links *Arhgap11a*-mediated RGC morphology to neuronal migration.

Given these defects in radial migration, we next asked if acute *Arhgap11a* loss from RGCs affects cell composition and cortical lamination.³⁹ We assessed brains that had been electroporated at E15.5 with either scrambled siRNAs or *Arhgap11a* siRNAs. A 2-day knockdown of *Arhgap11a* had no striking impact on progenitor proliferation; indeed, 40% of control and mutant IUE'd cells were Ki67 positive, and the RGC (Sox2+) and intermediate precursor (IP) (Tbr2+) fractions were normal (Figures S3A–S3D). Consistent with this, at P0, we noted no significant difference in the distribution of electroporated cells or in the density of cells between conditions (Figures S3E–S3H). Although we cannot rule out earlier roles in cell fate specification, these data suggest that acute *Arhgap11a* knockdown does not grossly influence neurogenesis.

We next assessed whether the radial migration deficiency was associated with altered laminar organization in the P0 cortex (Figure S4A). We first quantified the distribution of neurons born several days prior to *Arhgap11a* siRNA treatment (layers V and VI) (Figure S4A). Tbr1+ layer VI neuron distribution was similar between knockdown and control mice (Figures S4B and S4C). In comparison, Ctip2 layer V neurons showed a slight but significant altered distribution (Figures S4D and S4E). This is consistent with the expectation that, at the time of acute RGC knockdown, virtually all layer VI neurons had completed their migration while some layer V neurons were still migrating. We then examined the distribution of neurons with peak genesis 1 day prior to, or simultaneous with electroporation (layers IV and II/III). Compared with scrambled control, fewer ROR-β+ layer IV neurons localized to superficial layers in *Arhgap11a* knockdown brains (Figures S4F and S4G). Lhx2+ layer II/III neurons were also altered in their distribution (Figures S4H and S4I). Notably, the distribution of Ctip2+ and ROR-β+ neurons in P21 *Arhgap11a*-deficient cortices was also significantly altered, particularly layer IV neurons (Figures S4J–S4N). This phenotype is striking given that it resulted from an acute embryonic knockdown about 25 days earlier. Altogether, these results indicate that acute *Arhgap11a* knockdown in RGCs cause subtle but significant impairments of the laminar organization of cortical neurons at both P0 and P21. As *Arhgap11a* is not expressed in excitatory neurons (Figures S1B–S1F), this suggests that neuronal organization is influenced by the non-cell-autonomous requirements of *Arhgap11a* in the RGC scaffold.

***Arhgap11a* is essential for RGC basal endfeet morphology and positioning of cortical interneurons**

RGC basal process morphology is dynamic and complex not only in the IZ but also near the pial surface in the marginal zone (MZ).^{15,16} Indeed, morphological analyses of RGCs show that endfeet number is constant through E14.5 and increases by E16.5.¹⁶ Moreover, consistent with others' findings¹⁵ we

observed by live imaging of E16.5 embryonic brain slices, that basal processes can be highly dynamic in the MZ (Video S3). ARHGAP11A localization to distal regions of RGC basal processes and endfeet coincides developmentally with this RGC dynamic behavior and endfeet complexity¹⁶ (Figures 1C–1H). Therefore, we tested whether *Arhgap11a* regulates basal process complexity in the MZ.

Toward this, we employed 3D reconstructions of either control or siRNA-treated membrane-bound-EGFP+ basal processes at the pia, introduced by IUE at E15.5. We specifically quantified the RGC arbor and branching of basal processes (>5 μm long⁴⁰) in the MZ (Figures 3A–3C). In RGCs transfected with scrambled siRNAs, we observed extensive branches emanating from the basal process in the MZ, as described previously (Figures 3C–3E).^{15,16} In contrast, *Arhgap11a* siRNA knockdown induced a significant decrease in both branching complexity and the total number of branches. This effect persisted after 48 h, albeit to a lesser extent (Figures S5A–S5D). To orthogonally validate this phenotype, we used *Arhgap11a* germline knockout (KO) mice (Figures S5F–S5H). Analysis of endfeet of the E16.5 KO brains showed significant reductions in complexity, paralleling that seen in the siRNA-depleted brains (Figures S5I–S5K). Given these mice are germline deficient in *Arhgap11a*, we opted to use siRNA-treated brains for further analysis of acute phenotypes.

We next evaluated whether *Arhgap11a* influences the basal endfoot area contacting the basal lamina. To do this, we generated endfoot preparations from electroporated brains, which included meninges, basal lamina, and RGC basal endfeet.²¹ Imaging basal endfeet with an en-face view, we quantified an average basement membrane coverage of ~9 μm² by control RGC endfeet (Figures 3F and 3G). However, this coverage significantly diminished by 25% in RGC endfeet depleted of *Arhgap11a*. Of note, there was no observable impact of the *Arhgap11a* knockdown on the number of endfeet per RGC (Figure S5E). Altogether, these results show that *Arhgap11a* is essential for RGC morphology in the MZ.

This raises the question of what is the impact of these morphology defects upon cortical architecture at the pia? To first assess this question, we used serial-blockface electron microscopy (SBFEM) to visualize RGC basal processes and endfeet structure as well as their interactions with the surrounding MZ niche (Figures 3H–3M). 3D reconstructions of RGCs highlighted that basal endfeet tile the basement membrane, forming a tight interface with the basement membrane (Figures 3H and 3I). RGC basal processes and endfeet also directly contacted surrounding cells and neurites of the MZ. For example, we reconstructed tangentially directed cells (presumably interneurons or Cajal-Retzius neurons) contacting a multitude of RGC basal processes and branches (Figures 3L and 3M).

Given that *Arhgap11a* controls RGC morphology in the MZ, we asked whether depleting *Arhgap11a* in RGCs could non-cell autonomously affect resident cells of the MZ, including Cajal-Retzius neurons and interneurons.^{41–47} We introduced scrambled or *Arhgap11a* siRNAs into RGCs using IUE at E15.5 and analyzed cell density in the MZ 24 h later, focusing on the region containing EGFP+ RGC basal processes and endfeet (Figures 4A and 4B). Strikingly, compared with the control,

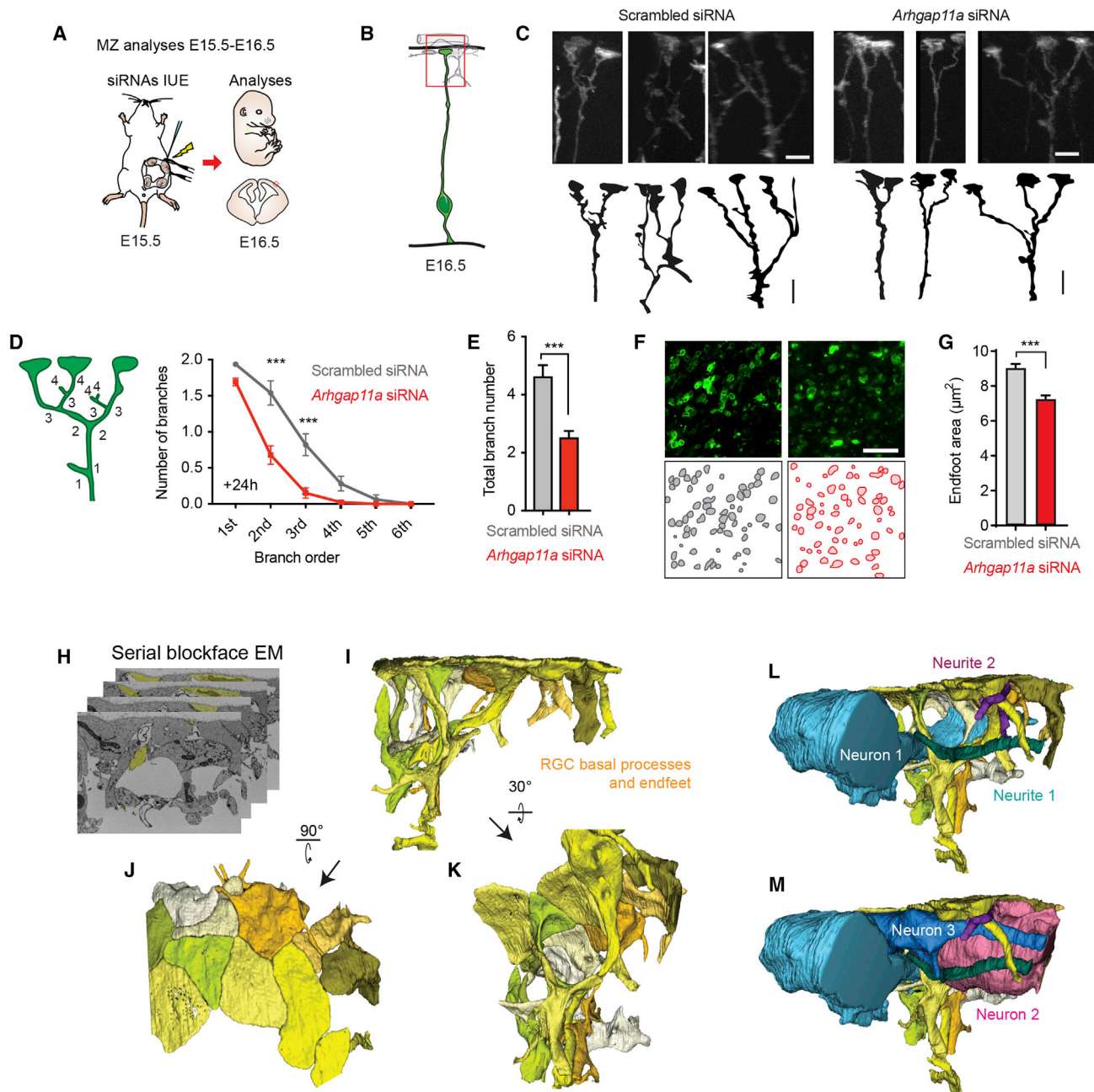


Figure 3. *Arhgap11a* promotes RGC basal process and endfeet complexity and interneuron numbers in the marginal zone

(A) Schematic overview of the experiments in (B)–(F) which examine acute impact of RGC knockdown upon RGC basal process and endfeet in the marginal zone (MZ).

(B) Region analyzed in the experiments.

(C) Representative images showing basal process and endfeet complexity in the MZ in IUE'd RGCs. Tracing of images is below.

(D and E) Method to define branch orders in RGC basal processes in MZ (D, left) and quantification of branch complexity (D, right), and average total branch number per RGC (E) (scrambled: $n = 78$ cells, 6 brains, 4 independent experiments, *Arhgap11a*: $n = 78$ cells, 5 brains, 4 independent experiments, two-way ANOVA, Sidak post hoc analyses to compare branch order).

(F and G) Representative images (F) for quantification of endfoot-basal lamina contact area in the MZ (G) (scrambled: $n = 351$ endfeet, 6 brains, 4 independent experiments, *Arhgap11a*: $n = 351$ endfeet, 6 brains, 4 independent experiments, Mann-Whitney test).

(H–M) 3D reconstructions of the MZ niche using serial-blockface electronic microscopy shows tight interactions between presumed (blue, pink) interneurons, Cajal-Retzius neurons, and RGC basal processes and endfeet (yellow).

siRNAs, small interfering RNAs; IUE, *in utero* electroporation; MZ, marginal zone; EM, electronic microscopy. * p value < 0.05. *** p value < 0.001. Individual data points represent different brains. Scale bars: (C) 5 µm, (F) 15 µm. Graph and bar plots, means ± SEM.

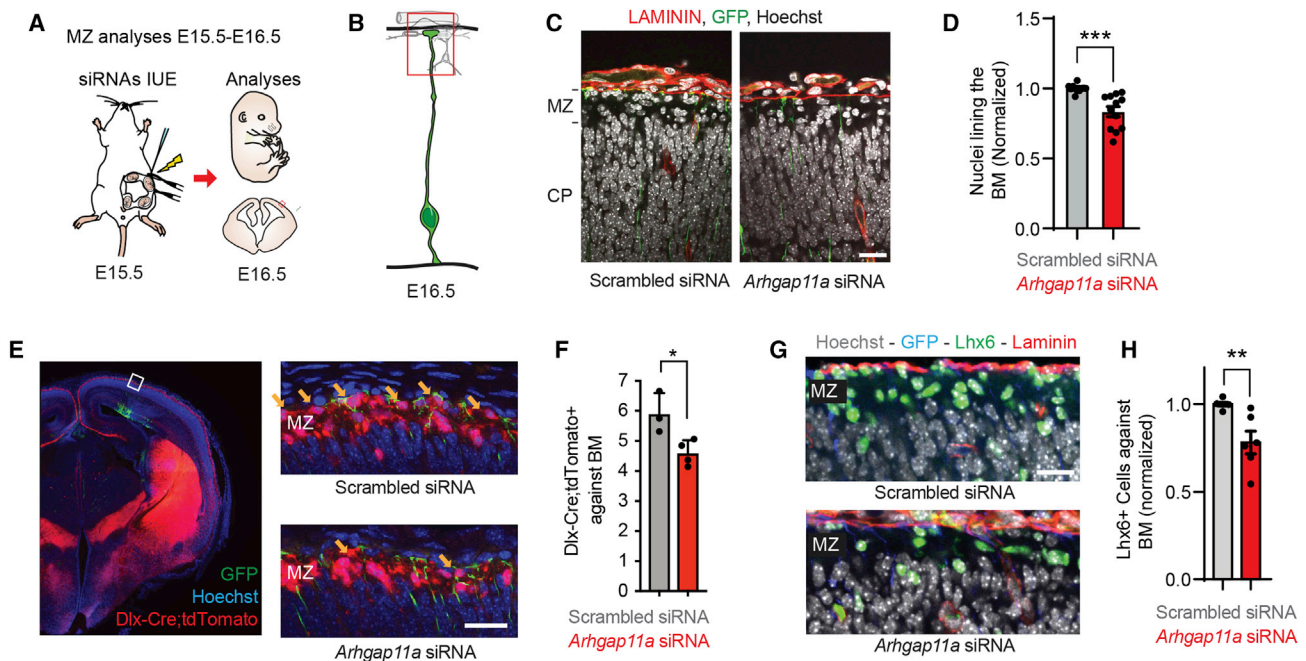


Figure 4. *Arhgap11a* influences interneuron positioning in the marginal zone

(A) Schematic overview of the experiments in (B)–(H) which examine acute impact of RGC knockdown upon interneuron positioning in the marginal zone (MZ). (B) Region analyzed in the experiments. (C) Immunofluorescence depicting nuclei (white, Hoechst) and laminin (red) in GFP (green) electroporated regions. (D) Quantification of the number of nuclei lining the basement membrane (BM), across a region of analysis (scrambled: n = 9 brains, 5 independent experiments, *Arhgap11a*: n = 4 brains, 4 independent experiments, unpaired t test). (E) Immunofluorescence depicting Hoechst+ nuclei (blue), tdTomato+ interneurons (red) in GFP (green) electroporated region, with higher magnification images on right. Yellow arrows, tdTomato+ interneurons located against the basement membrane. (F) Quantification of Tomato+ interneurons lining the BM in indicated brains (scrambled: n = 3 brains, 1 experiment, *Arhgap11a*: n = 4 brains, 1 experiment, unpaired t test). (G) Immunofluorescence depicting Hoechst+ nuclei (blue), Laminin (red), Lhx6+ interneurons (green) in GFP (blue) electroporated region. (H) Quantification of Lhx6+ interneurons lining the BM in indicated brains (scrambled: n = 6 brains, 4 experiments, *Arhgap11a*: n = 6 brains, 3 experiments, Mann-Whitney test). siRNAs, small interfering RNAs; IUE, *in utero* electroporation; MZ, marginal zone; EM, electronic microscopy. *p value < 0.05. ***p value < 0.001. Individual data points represent different brains. Scale bars: (C and E) 25 μ m, (G) 20 μ m. Bar plots, means \pm SEM.

Arhgap11a knockdown in RGCs significantly reduced the number of Hoechst+ cells lining the LAMININ+ basement membrane (Figures 4C and 4D).

We then quantified interneurons, which migrate tangentially from the ganglionic eminences into the cortex via 3 streams, including along the MZ. Interneurons that migrate through the MZ invade the CP, forming axons that extend from layer I along the way.⁴⁸ This interneuron population is thus critical for cortical circuitry. We employed IUE of siRNAs and EGFP in *Dlx-Cre; Ai14* animals, in which interneurons express tdTomato (Figures 4E and 4F). Strikingly, *Arhgap11a* knockdown in RGCs significantly decreased the number of interneurons lining the basement membrane, where endfeet are located (Figure 4F). Consistent with this, we quantified fewer Lhx6+ interneurons lining the basement membrane following acute *Arhgap11a* knockdown (Figures 4G and 4H). In contrast, there was no obvious alteration in Calretinin+ cells, which label small numbers of interneurons but at this stage mainly label Cajal-Retzius neurons⁴² (Figures S5L and S5M). These results demonstrate that *Arhgap11a* acts non-cell autonomously within RGCs to promote

the position of a population of interneurons in the MZ. Taken together with the discovery that basal process complexity is associated with excitatory neuron migration (Figures 2, S2, and S3), these findings broadly link RGC morphology in basal structures to non-cell autonomous positioning of neurons.

***Arhgap11a* mRNA is actively transported to RGC basal endfeet via a 5' UTR *cis* element**

Our data indicate that *Arhgap11a* is essential for RGC basal morphology and neuronal migration. This begs the question of how *Arhgap11a* controls these RGC subcellular compartments, and specifically the basal endfeet architecture. Notably, both *Arhgap11a* mRNA and protein are subcellularly enriched and co-localize in RGC endfeet (Figure 1), as seen for other transcripts.²¹ This suggests that RNA transport and local translation of *Arhgap11a* may enable rapid and specific control of RGC basal endfoot morphology.

Active mRNA transport depends on *cis*-localization elements within mRNAs.⁴⁹ Therefore, we first determined the sequence(s) within *Arhgap11a* mRNA that confers endfoot localization. To

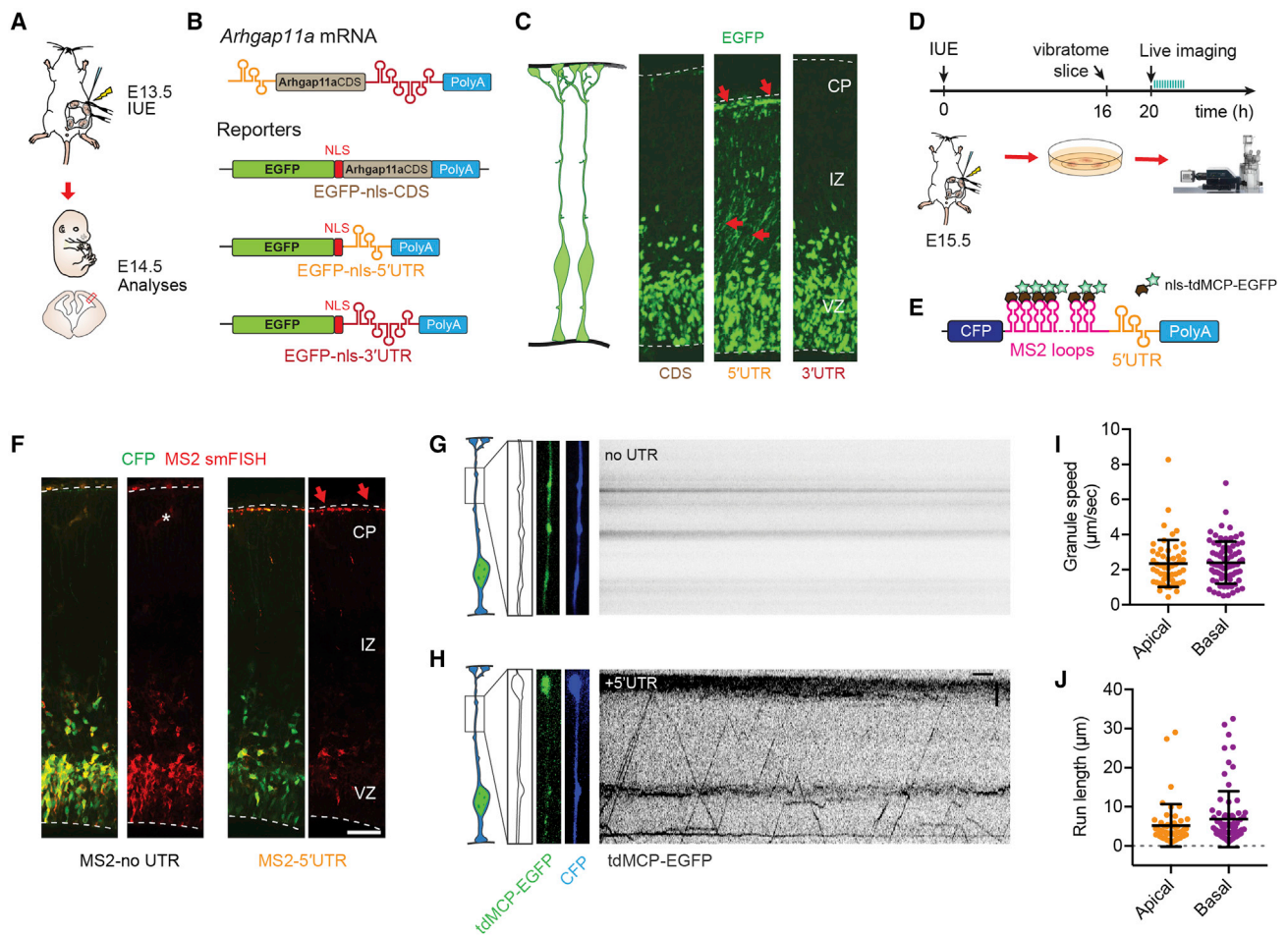


Figure 5. *Arhgap11a* mRNA is actively transported to radial glial basal endfeet via a 5' UTR element

(A and B) Schematic overview (A) of the strategy used in (B and C) to determine the endfoot localization element in *Arhgap11a* mRNA using a 1-day electroporation of indicated reporter constructs (B).

(C) EGFP-nls localizes to RGC basal endfeet only when the *Arhgap11a* 5' UTR is present, but not in CDS alone or containing 3' UTR.

(D and E) Schematic overview (D) of the strategy used in (F)–(J) to visualize transport of *Arhgap11a* mRNA reporters (E) in RGC basal processes.

(F) smFISH (red) targeting MS2 stem-loop RNA sequences shows *Arhgap11a* 5' UTR induces RNA localization from cell bodies to RGC basal process and endfeet (CFP, green).

(G and H) Kymographs showing absence (G) and presence (H) of MS2-tagged mRNA transport in RGC basal process over a 1-min period, in no UTR and 5' UTR, respectively.

(I and J) Quantification of similar average speeds of MS2-tagged mRNA transport in RGC basal processes (I) and average run lengths (J) in both apical and basally directed movements. n = 126 EGFP+ punctae, 11 cells, 2 brains, 2 independent experiments.

IUE, *in utero* electroporation; CDS, coding sequence; UTR, untranslated region; nls, nuclear localization signal; tdMCP, tandem MS2-coat protein; CFP, cyan fluorescent protein; CP, cortical plate; IZ, intermediate zone; VZ, ventricular zone. Scale bars: (F) 50 μ m, (G and H) horizontal axis: 5 s, vertical axis: 5 μ m. Graphs, mean \pm SEM.

this end, we generated three reporters containing EGFP with a nuclear localization sequence (NLS-EGFP), together with either the *Arhgap11a* (1) 5' UTR (527 bp), (2) coding sequence (2,942 bp), or (3) 3' UTR (1,057 bp). To test for position-independent localization capacity, these elements were included at the 3' end of the reporter, followed by a poly(A) sequence (Figures 5A and 5B). The reporters were introduced by IUE at E13.5 and E14.5 brains were collected. As previously shown,²² EGFP reporters that are not transported are nuclear, whereas EGFP localization in the basal process and endfeet indicates transport from the RGC cell body and translation in the basal

endfeet. Using this strategy, we discovered that the endfoot RNA localization sequence of murine *Arhgap11a* mRNA resides in its 5' UTR and not in its coding or 3' UTR sequences (Figures 5A–5C). This was further confirmed by *in situ* hybridization targeting the EGFP mRNAs in electroporated brains (Figures S6A and S6B).

Having determined the sequence element of *Arhgap11a* sufficient for its localization to endfeet, we next asked if *Arhgap11a* is actively transported to endfeet. To visualize *Arhgap11a* mRNA transport, we live imaged mRNAs in RGC basal processes within organotypic brain slices as previously.²¹ This technique uses a

reporter plasmid with CFP and a 3' UTR containing 24 MS2 stem loops, and a second plasmid containing the nuclear localized tdMCP-EGFP protein with high affinity for RNA MS2 stem loops (Figures 5D and 5E).⁵⁰ Co-transfecting both plasmids allows the indirect visualization of mRNAs. Two different reporters encoding CFP were generated: one control sequence with MS2 loops only (MS2-no UTR) and the *Arhgap11a* 5' UTR sequence downstream of the MS2 loops (MS2-5' UTR). These reporters were transfected into RGCs by IUE at E15.5, and organotypic slices were generated at E16.5. In brains electroporated with either control MS2-noUTR or the MS2-5' UTR, we observed diffuse EGFP+ in the nucleus and cytoplasm of CFP+ RGCs, demonstrating the efficiency of the technique (Figure 5F). smFISH targeting MS2 loops in fixed electroporated brain sections corroborated the localization of the 5' UTR reporter to basal endfeet.

Having established these tools, we then performed live imaging of electroporated EGFP+ mRNAs in the basal process of E16.5 RGCs within organotypic brain slices. This allowed us to visualize active directed transport of MS2-5' UTR EGFP+ mRNAs (Figures 5D, 5E, 5G, and 5H; Video S4). Kymograph analyses highlighted bidirectional movements, with frequent changes of orientation in the 5' UTR reporter but not the no UTR (Figures 5G and 5H). 60% of the observed movements were directed toward the basal endfeet (Figure S6C). We measured average speeds of 2.5 $\mu\text{m/s}$ and tracked single uninterrupted movements of up to 32 μm , with an average of 6 μm (Figures 5I and 5J). These speeds and processivity are consistent with active microtubule-dependent transport, and similar to those observed previously for other mRNA reporters.²¹ Altogether, these results demonstrate that *Arhgap11a* mRNA is actively transported to basal endfeet, arguing that its subcellular localization may ultimately impact basal endfeet.

ARHGAP11A subcellular localization in RGC basal processes and endfeet relies upon local translation in the basal endfeet

The presence of ARHGAP11A protein in the basal process and endfeet could be explained by at least two possibilities: synthesis of ARHGAP11A in the soma followed by protein transport into the basal process and endfeet, and/or local production of ARHGAP11A within endfeet. Given the active transport of *Arhgap11a* mRNA along the basal process and enrichment in endfeet, we predicted the latter. To test this directly, we generated reporters expressing ARHGAP11A full-length protein fused to the fluorescent photoconvertible protein DENDRA2. Two reporters were generated: a control producing an mRNA devoid of any localization sequence (DENDRA2-no UTR) and another including the 5' UTR of *Arhgap11a* sufficient for RNA localization at the endfoot (DENDRA2-5' UTR, Figure 6A). DENDRA2 fluorescent signal was analyzed in fixed E16.5 brains after IUE at E15.5. smRNA FISH targeting the *Dendra* sequence confirmed the transport of the 5' UTR reporter to basal endfeet (Figures S7A–S7C). Expression of either the control or 5' UTR reporter resulted in a DENDRA2 signal in RGC cell bodies (Figures 6B and 6C), recapitulating endogenous protein expression (Figures 1C–1H). However, the DENDRA2 signal in the RGC basal endfeet was only evident in brains electroporated with the 5' UTR reporter (Figures 6B–6E). In these brains, ARHGAP11A localization was

also evident along the basal processes within the IZ and CP (Figures 6D and 6E). This localization was not context dependent, as placement of the 5' UTR in either the 5' or 3' part of the reporter led to similar enriched localization to the basal endfeet (Figure S7D). This strongly argues that ARHGAP11A protein localization within endfeet and along the basal process relies upon subcellular localization of the transcript.

Next, we used live imaging to directly test if ARHGAP11A can be locally synthesized within endfeet. Using the DENDRA2 constructs described above together with CFP to identify electroporated endfeet, we employed live imaging using *ex vivo* endfoot preparations, consisting of isolated basal endfeet connected to the basement membrane and overlying meninges, as previously described^{21,51} (Figure 6F). UV-exposure was used to photoconvert the DENDRA2 signal in isolated endfeet from green to red (Figures 6G and 6H). Subsequently, the recovery of green fluorescence was monitored over 45 min. Since basal endfeet are completely disconnected from the cell body, any new green fluorescence is a robust readout of *de novo* local protein synthesis. No recovery was noted in the negative control lacking the 5' UTR (Figure 6G and 6J). In contrast, green fluorescence increased in endfeet expressing *Dendra2*-5' UTR over 45 min (Figures 6H–J). Importantly, these kinetics were completely abolished in the presence of the translation inhibitor anisomycin (Figure 6J). Altogether, these data demonstrate that active *Arhgap11a* mRNA transport and local translation in endfeet enable subcellular expression of ARHGAP11A protein in RGC basal structures.

Locally synthesized ARHGAP11A controls basal process morphology through RhoGAP activity which influences interneuron positioning

ARHGAP11A mRNA and protein localize to RGC basal endfeet coincident with increasingly complex endfoot morphology. Given this, and the observation that mRNA is actively transported and locally translated during development, we postulated that local synthesis of this RhoGAP controls endfoot morphology and thus impacts interneuron positioning along the basal membrane. To test this, we used IUE at E15.5 to introduce *Arhgap11a* siRNAs together with *Arhgap11a* full-length protein either with or without the 5' UTR (Figures 7A–7D). Importantly, both *Arhgap11a* cDNAs were resistant to the transfected siRNAs. Compared to siRNA knockdown alone, co-expression of siRNAs with full-length *Arhgap11a* did not restore RGC basal process complexity in the MZ (Figures 7E–7G, 7J, and 7K). In contrast, the introduction of *Arhgap11a*-5' UTR completely restored the RGC morphology phenotype (Figures 7E, 7F, 7H, 7J, and 7K). Further, while non-localized *Arhgap11a* failed to rescue the reduced endfoot area, localized ARHGAP11A did (Figures 7L and 7M). Of note, expression of *Arhgap11a* with or without the 5' UTR alone, but without siRNA, did not affect RGC branching complexity in the MZ, indicating there is no overt overexpression phenotype (Figures S8A–S8C). These findings demonstrate that subcellular localization of *Arhgap11a* via its 5' UTR is critical for proper basal process morphology.

Is RGC morphology dependent upon ARHGAP11A Rho GTPase function? To address this question, we generated a previously described GAP-deficient (GD) form of ARHGAP11A.²⁵ This GD-*Arhgap11a*-5' UTR reporter contains a single missense

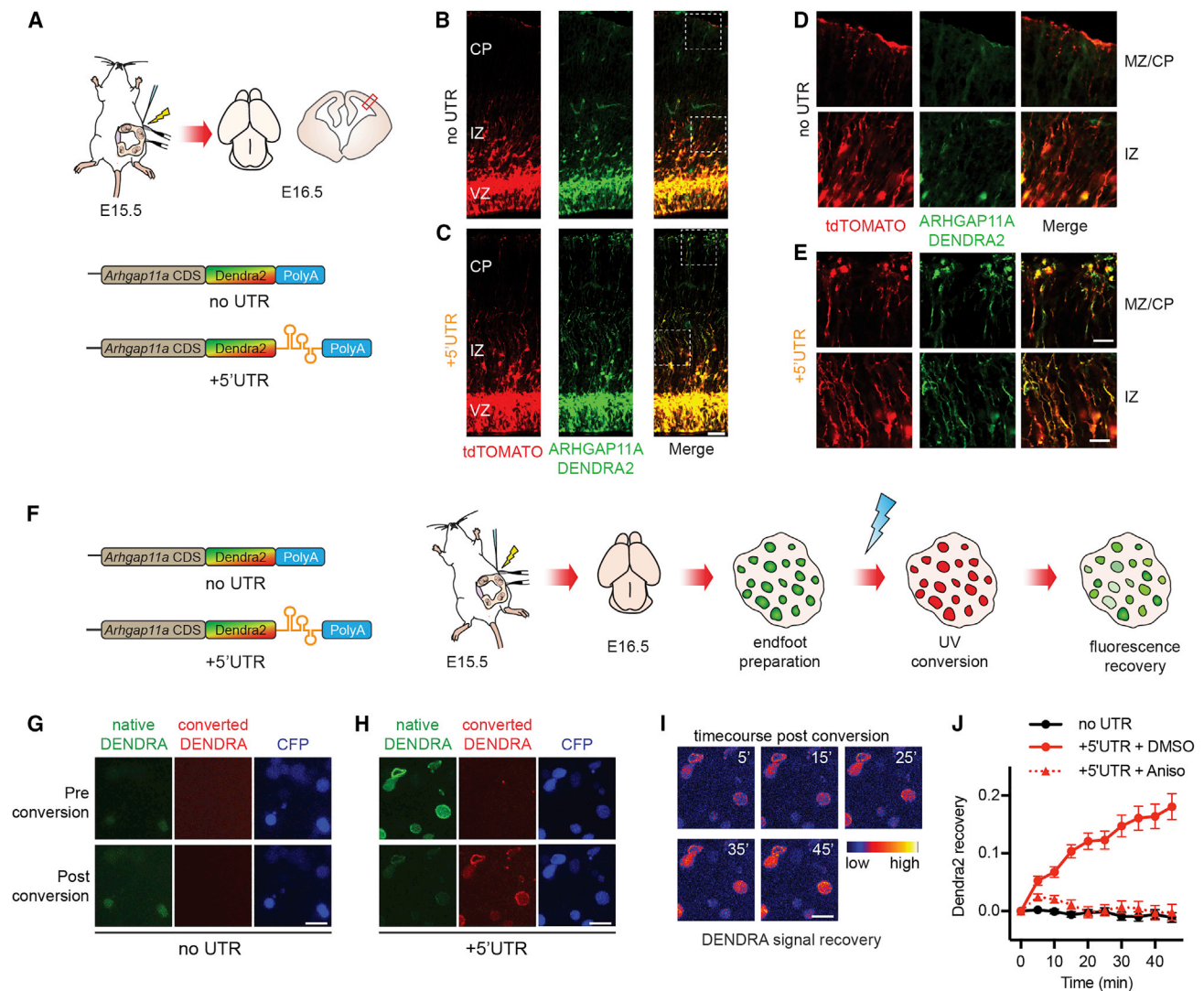


Figure 6. ARHGAP11A protein localization to RGC basal processes and endfeet relies on local translation of *Arhgap11a* mRNA in basal endfeet

(A) Schematic overview of the strategy used in (B)–(E) to test if *Arhgap11a* mRNA localization mediates ARHGAP11A expression in RGC basal endfeet and basal processes.

(B–E) Immunofluorescence of tdTomato electroporated RGCs (red) and ARHGAP11A fusion reporter (green) containing no UTR (top) or 5' UTR (bottom). High magnification images (D and E) reflect ARHGAP11A protein localizes to RGC basal endfeet (MZ/CP) and basal processes (IZ) only with *Arhgap11a* 5' UTR.

(F) Schematic overview of the strategy used in (H)–(K) to visualize local translation of *Arhgap11a* in RGC basal endfeet.

(G and H) Images showing ARHGAP11A-DENDRA fluorescence in RGC basal endfeet pre (top) and post- (bottom) photoconversion in no UTR (G) or 5' UTR (H) conditions.

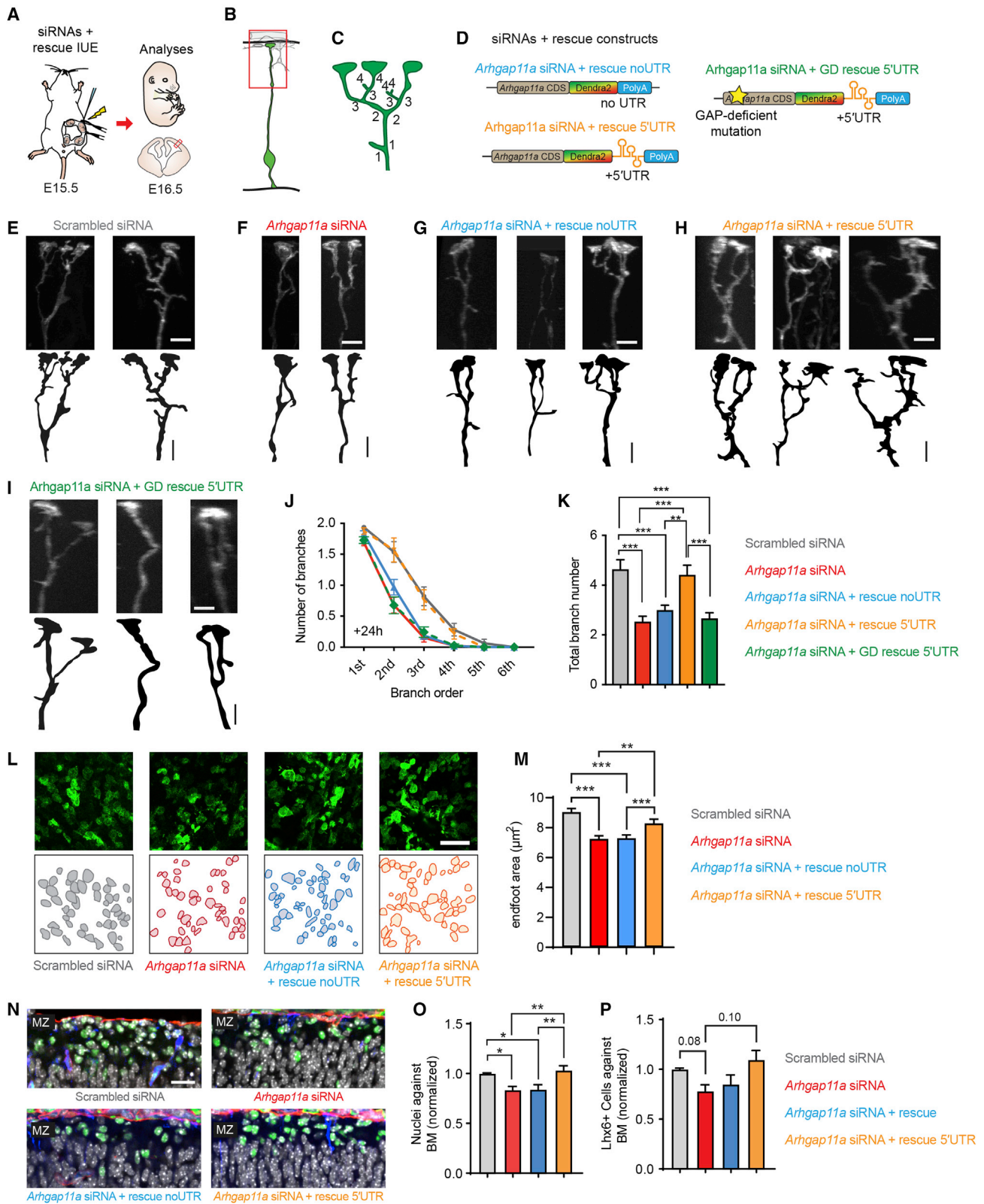
(I) Time course showing recovery of native DENDRA signal in the +5' UTR condition, as pseudocolored using indicated scale (time, min).

(J) Quantification of positive recovery of native DENDRA signal in RGC basal endfeet only with 5' UTR +DMSO (red, solid line) relative to both the no UTR condition (black) and anisomycin treatment (red, dotted line). no UTR: n = 27 endfeet, 2 brains, 2 independent experiments, 5' UTR + DMSO: n = 62 endfeet, 3 brains, 3 independent experiments, n = 69 endfeet, 3 brains, 3 independent experiments, two-way ANOVA interaction time × condition: p value < 0.0001.

UTR, untranslated region; CDS, coding sequence; Aniso, anisomycin. Scale bars: (B and C) 100 μm; (D and E) 20 μm; (G–I) 5 μm. Graph, means ± SEM.

mutation, R87A (Figure 7D). The transcript encoded by this mutant ARHGAP11A localized properly to endfeet (Figure S7C). However, like the no UTR construct, it also failed to rescue the siRNA-mediated basal process phenotype in the MZ (Figures 7I–7K). This indicates local Rho GAP activity mediates RGC basal process complexity in the cortex.

Acute depletion of *Arhgap11a* from RGCs impaired their basal morphology and led to significantly fewer Hoechst+ cells and interneurons at the basement membrane (Figures 3 and 4). Given this, we asked whether rescuing RGC morphology at the pia was sufficient to also recover interneuron positioning. Re-introduction of full-length *Arhgap11a*, which lacked a localization



(legend on next page)

element, failed to rescue both nuclei number and interneuron number (Figures 7N–7P). This is consistent with the inability of non-localized *Arhgap11a* to rescue endfoot complexity and area (Figures 7K and 7M). In contrast, co-expression of *Arhgap11a*-5' UTR with the siRNAs recovered nuclei as well as interneuron numbers. Together, these results demonstrate that RGCs employ mRNA transport and local protein synthesis of *Arhgap11a* at RGC endfeet to fine-tune basal process morphology and direct neurons to their proper position (Figure 8).

DISCUSSION

By controlling neurogenesis and neuronal position, RGCs are crucial for orchestrating the development of the cerebral cortex. Yet how the polarized morphology of RGCs dictates these functions is poorly understood. Here, we demonstrate that RGCs and cortical architecture rely upon exquisite temporal and spatial control of gene expression via active mRNA transport and local translation. We show that RhoA-GAP activity is spatially and acutely activated via local translation in RGC basal endfeet to control their morphology and non-cell autonomously influence interneuron positioning. Our study demonstrates that mRNA localization and local translation in RGCs are essential for brain development *in vivo*. This implicates a dynamic new gene regulatory mechanism by which progenitors shape brain development across species.

mRNA transport and local translation control RGC morphology

Using live and fixed *ex vivo* and *in vivo* imaging, we demonstrate that *Arhgap11a* mRNA undergoes active transport, subcellular localization, and local translation in RGC basal endfeet. This discovery adds *Arhgap11a* to a short but growing list of over 100 transcripts that we and others have shown are subcellularly localized to RGC endfeet.^{21,22,52,53} This further reinforces the finding that RGC basal processes are highways for active

mRNA transport and exhibit subcellular translation at the pia.^{51,54}

To date, the functions of any subcellularly localized transcript in endfeet are unknown. We address this significant gap by demonstrating that at least one function of subcellular RNA localization in RGCs is to mediate local morphology. This is based on several lines of evidence. First, *Arhgap11a* mRNA and protein localization to basal endfeet and basal processes relies upon its 5' UTR and coincides with critical developmental stages when RGC complexity increases.¹⁶ Second, acute *Arhgap11a* depletion decreased RGC complexity in basal structures; importantly and strikingly, this phenotype was rescued only upon expression of endfoot-localized, functional *Arhgap11a*. Hence, RGC basal structures rely upon coordinated active mRNA transport, local translation, and local and acute Rho GTPase function.

Why do RGCs require protein production in endfeet rather than transport existing proteins from the cell body? Translation of an average protein takes about 1 min, whereas transcription is 10 times longer.⁵⁵ Thus, local expression is both highly efficient and energetically favorable, enabling rapid changes in RGC morphology as the brain grows radially and tangentially. Our previous data show that endfeet contain over 100 significantly enriched FMRP-bound mRNAs, including those encoding cytoskeletal and signaling regulators,²¹ suggesting this may be a widespread mechanism in RGCs. While not significantly enriched in this dataset, *Arhgap11a* could be a low-affinity FMRP target. Migrating fibroblasts and neuronal growth cones and spines also contain similar classes of localized transcripts,^{54,56–58} suggesting our findings may extend to other cell types. Beyond morphology, localized mRNAs in endfeet may also have additional functions, such as promoting cell fates or intra- and extracellular signaling.²²

How does ARHGAP11A protein synthesized in endfeet influence complexity along the basal process? Our data suggest that endfeet ARHGAP11A moves into the basal process either passively, via diffusion, or actively. Indeed, exogenous ARHGAP11A reporter localization to the basal process relied

Figure 7. Locally synthesized ARHGAP11A controls basal process morphology through GAP activity

(A) Schematic overview of the strategy used in (B)–(K) to assess rescue of RGC endfeet morphology.

(B and C) Method to define branch orders in RGC basal processes in MZ.

(D) Rescue constructs used in experiments.

(E–I) Representative images showing basal process complexity at the level of the MZ in RGCs treated as indicated.

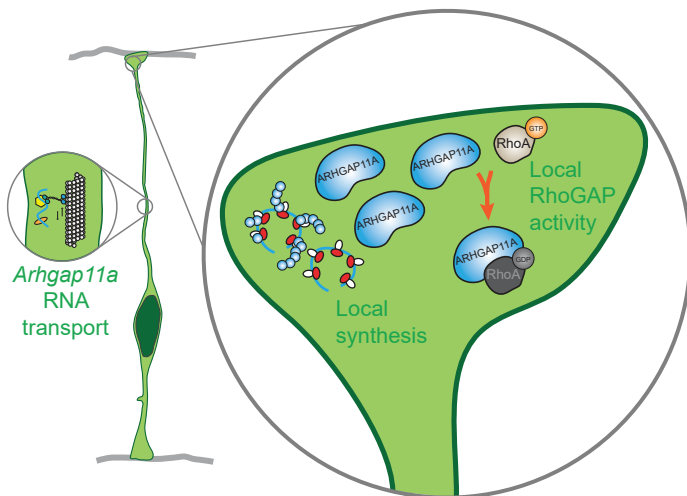
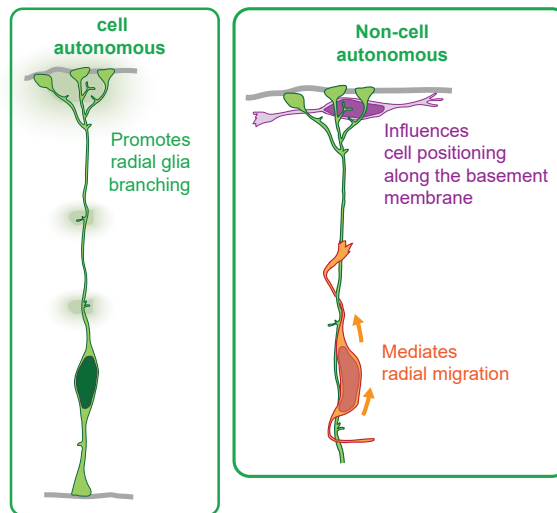
(J and K) Quantification of basal process and endfoot complexity at the level of the MZ in RGCs (scrambled: n = 78 cells, 6 brains, 4 independent experiments, *Arhgap11a*: n = 78 cells, 5 brains, 4 independent experiments, *Arhgap11a* + rescue no UTR: n = 97 cells, 6 brains, 4 independent experiments, *Arhgap11a* + rescue 5' UTR: n = 56 cells, 4 brains, 3 independent experiments, *Arhgap11a* + GD rescue 5' UTR: n = 78 cells, 4 brains, 2 independent experiments, (J) two-way ANOVA: p value < 0.0001, (K) one-way ANOVA: p < 0.0001, Tukey's post hoc comparisons).

(L and M) Quantification of endfoot-basal lamina contact area in RGCs (scrambled: n = 351 endfeet, 6 brains, 4 independent experiments, *Arhgap11a*: n = 351 endfeet, 6 brains, 4 independent experiments, *Arhgap11a* + rescue no UTR: n = 454 endfeet, 7 brains, 4 independent experiments, *Arhgap11a* + rescue 5' UTR: n = 200 endfeet, 4 brains, two independent experiments, Kruskal-Wallis test).

(N) Images of interneurons (LHX6, green) along the basement membrane (BM, red) in different genotypes.

(O and P) Quantification of the positioning of DAPI cells (white) and interneurons along the basement membrane (nuclei against BM: scrambled: n = 13 brains, 9 independent experiments, *Arhgap11a*: n = 12 brains, 7 independent experiments, *Arhgap11a* + rescue no UTR: n = 14 brains, 8 independent experiments, *Arhgap11a* + rescue 5' UTR: n = 13 brains, 8 independent experiments, one-way ANOVA followed by Tukey post hoc analyses; Lhx6+ cells against the BM: scrambled: n = 6 brains, 4 independent experiments, *Arhgap11a*: n = 6 brains, 3 independent experiments, *Arhgap11a* + rescue: n = 6 brains, 3 independent experiments, *Arhgap11a* + rescue 5' UTR: n = 8 brains, 4 independent experiments, Brown-Forsythe and Welch ANOVA followed by Dunnett T3 post hoc analyses).

IUE, *in utero* electroporation; siRNA, small interfering RNA; CDS, coding sequence; UTR, untranslated region; GD, Rho-gap-deficient. ** p value < 0.01. *** p value < 0.001. Scale bars, (E–I) 5 μ m; (L) 15 μ m; (N) 20 μ m. Graphs, means \pm SEM.

Arhgap11a mRNA transport and local translation**Arhgap11a functions****Figure 8. Model for major findings of this study**

Arhgap11a mRNA is actively transported in RGC basal process to basal endfeet. In basal endfeet local synthesis of ARHGAP11A protein enables expression in basal structures and local RhoGAP activity, thus promoting radial glia branching and interneuron position. *Arhgap11a* is non-cell autonomously required in RGCs for migration of excitatory neurons and positioning of inhibitory neurons.

entirely on an endfoot localization element. We also observed sporadic enrichment of *Arhgap11a* mRNAs along the basal process, which could result from collective RNA trafficking to basal endfeet. Further, such enrichment could indicate local translation “hotspots” in the basal process, as proposed in axons.⁵⁹

Acute, spatial regulation of a Rho-GTPase modulator controls RGC morphology and neuronal positioning

We demonstrate that localized, acute expression of GTPase regulators mediates subcellular architecture. GTPases, including RhoA, can promote filopodial activity and morphology in RGC basal structures, ultimately influencing neuronal migration and neurogenesis.^{12,15,23,60–62} RhoA signaling via the microtubule regulator Memo has also been linked to the tiled distribution of RGC basal processes, hyperbranching, and migration defects.^{38,63} Consistent with these roles, *Rho* KO mice have heterotopia.²³ In contrast, consistent with the prediction that *Arhgap11a* loss induces excessive Rho signaling, *Arhgap11a* depletion reduces branching and impairs radial migration. Although acute knockdown did not drastically impact neurogenesis, we cannot rule out roles for *Arhgap11a* in cell fate at other stages. Thus, our findings are generally consistent with known Rho functions. Our data provide a new mechanism to understand how ubiquitously expressed Rho GTPases are locally and rapidly controlled in polarized cells of the nervous system.

Taken together, this suggests that both simplified and excessive RGC branching impair the orientation of radially migrating neurons and ultimately influence lamination. We speculate that RGC basal process branches act as the “rungs of a ladder” for neurons. While the absence of rungs could impede neuronal movement along the basal process, overextended “rungs”

could be misinterpreted by neurons as basal processes, thus deviating them from their migratory path.

RGC morphology has been previously suggested to influence interneurons,⁴¹ however, our study is the first to formally demonstrate a functional link. Indeed, our data argue that RGC morphology and local ARHGAP11A function in the endfeet dictate interneuron positioning. This is reinforced by a recent study from our lab.⁶⁴ Interneurons traverse the developing cortex mainly via the VZ and MZ, and their ultimate destination is tied to their fate.⁶⁵ The population of migrating interneurons in the MZ is specialized, developing into Martinotti cells which leave axons in layer I as they invade the CP.⁴⁸ Of note, interneuron mis-positioning in the MZ was evident with just a 1-day acute depletion of *Arhgap11a* from RGCs, and perhaps more sustained manipulation of *Arhgap11a* in RGCs and thus the disruption of RGC morphology could ultimately impact cortical circuitry.

A key question is how ARHGAP11A, as well as endfeet complexity, influence interneuron positioning at the basement membrane. As suggested by 3D-EM data, cells closer to the basal lamina maintain a strong interface with basal endfeet. Endfeet may represent a “lawn” on which interneurons are “crawling” between. There may be direct cell-cell interactions between RGC endfeet and migrating interneurons that regulate their migration or promote their adhesion and restriction in the MZ, as well as short- or long-range signals. Our findings lay the groundwork for future studies using detailed live imaging and morphological studies to assess these possibilities. Of note, while we did not observe any positioning defects for Cajal-Retzius neurons, it is possible that some sub-types of CR neurons could be affected. However, our data align with the timing of Cajal-Retzius neuron migration, which is complete by E12.5, well before we acutely knocked down the expression of *Arhgap11a*.⁴⁶

Roles for subcellular RNA localization in human cortical development and disease

Given the conserved localization of *ARHGAP11A* in the human neocortex, we predict that this RhoGAP, and more broadly, mRNA localization, is critical in human RGCs. In human neocortices, the RGC basal process is significantly longer than in mice (up to several millimeters) and a prominent feature of the outer radial glia/basal radial glia (oRGs/bRGs).^{19,66,67} Further, in primates, lamellar expansions decorate the basal process and reside in close contact with migrating neurons.⁵ *ARHGAP11A* is highly expressed in both human RGCs and oRGs. Thus, localized pools of *ARHGAP11A* may promote human cortical development by influencing both RGC and oRG morphology.

During evolution, a hominid-specific partial duplication of *ARHGAP11A* led to the emergence of *ARHGAP11B*, which is linked to cerebral cortex expansion in humans. *ARHGAP11B* is also expressed in human neural progenitors, and its forced expression promotes progenitor proliferation and neuron production.^{67–69} In contrast to *ARHGAP11A*, *ARHGAP11B* lacks both GTPase activity and the RNA motif for localizing to RGC endfeet and indeed is not localized at the pia.³⁶ It is interesting to consider whether *ARHGAP11A* might be modulated by *ARHGAP11B* activity in the cell body of human RGCs, a relationship seen for other human-specific duplications like *SRGAP2*⁷⁰ or *NOTCH2NL*.^{71,72}

In sum, our findings establish novel and essential roles for local translation in maintaining RGC integrity. It is important to note that defects in RGC scaffolds underlie diverse neurodevelopmental diseases, including heterotopias and lissencephaly.^{13,73} This highlights the importance of investigating subcellular mRNA localization and local translation in RGCs toward understanding normal development and disease.

STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.neuron.2023.02.023>.

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AUTHOR CONTRIBUTIONS

L.-J.P. and D.L.S. conceived the idea and designed the study. L.-J.P. and D.L.S. wrote the manuscript. L.-J.P., J.L., K.J., Y.T., C.M.M., B.D., I.K.S., F.C.A., P.K., and S.S., performed experiments and analyses. P.V., F.P., and D.L.S. supervised and funded research. All authors reviewed, edited, and approved the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

INCLUSION AND DIVERSITY

We support inclusive, diverse, and equitable conduct of research.

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STAR METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Rabbit anti Arhgap11a	Bethyl	Cat#A303-097A, RRID: AB_10890540
Rabbit anti Arhgap11a	Abcam	Cat#ab113261, RRID: AB_10866587
Mouse anti Nestin	BD biosciences	Cat#556309, RRID: AB_396354
Rabbit anti Tbr1	Abcam	Cat#ab31940, RRID: AB_2200219
Rat anti Ctip2	Abcam	Cat#ab18465, RRID: AB_2064130
Mouse anti ROR-beta	R&D Systems	Cat#N7927, RRID: AB_1964364
Rabbit anti Laminin	Millipore	Cat#AB2034, RRID: AB_91209
Rabbit anti Calretinin	Swant	Cat#7697, RRID: AB_2721226
Rat anti Sox2	Thermo Fisher	Cat#14-9811-80, RRID: AB_11219070
Rabbit anti Tbr2	Abcam	Cat#ab183991, RRID: AB_2721040
Rabbit anti Ki67	Cell Signaling Technology	Cat#12202, RRID: AB_2620142
Experimental models: Organisms/strains		
Mouse: C57BL/6J	JAX	Cat#000664, RRID: IMSR_JAX:000664
Mouse: Dlx-Cre (Tg(dlx5a-cre) ¹ Mekk/J)	JAX	Cat#008199, RRID: IMSR_JAX:008199
Mouse: Arhgap11a ^{em1} (IMPC) ^J	KOMP	Cat#MGI:5907256, RRID: MMRRC_042323-JAX
Oligonucleotides		
siRNA: siRNA flex targetting <i>Arhgap11a</i>	Qiagen	Cat#SI00902111
siRNA: siRNA flex targetting <i>Arhgap11a</i>	Qiagen	Cat#SI00902132
siRNA: siRNA flex targetting <i>Arhgap11a</i>	Qiagen	Cat#SI00902118
siRNA: All Stars Negative Control	Qiagen	Cat#1027281
In situ hybridization probe target human <i>ARHGAP11A</i>	This paper	2117–2657 of human ARHGAP11A (NM_014783.5)
smFISH Probe Set: <i>Arhgap11a</i>	LGC Biosearch Technologies	Custom made
smFISH Probe Set: <i>MS2</i>	LGC Biosearch Technologies	Pilaz et al. ²¹
smFISH Probe Set: <i>Dendra</i>	LGC Biosearch Technologies	Pilaz et al. ²¹
qPCR primer: <i>Actb</i> FW	Thermo Fisher	AGATCAAGATCATTGCTCCT
qPCR primer: <i>Actb</i> Rev	Thermo Fisher	CCTGCTTGCTGATCCACATC
qPCR primer: <i>Pax6</i> FW	Thermo Fisher	TCTTTGCTTGGGAAATCCG
qPCR primer: <i>Pax6</i> Rev	Thermo Fisher	CTGCCGTTCAACATCCTTAG
qPCR primer: <i>Arhgap11a</i> FW	Thermo Fisher	GCAGGTGTGCCAAGGCGAAGT
qPCR primer: <i>Arhgap11a</i> Rev	Thermo Fisher	TGCAAGTCGCCAACCAACTTTCA
Taqman probe: <i>Tubb3</i>	Thermo Fisher	Cat#Mm00727586_s1, RRID#
Taqman probe: <i>Gapdh</i>	Thermo Fisher	<i>Gapdh</i>
Recombinant DNA		
Plasmid: pCAGGS-GFP	Gift from Dr. Nicholas Gaiano	Mizutani et al. ⁷⁴
Plasmid: pGLAST-EGFP-CAXX	Gift from Dr. Tarik Haydar	Gal et al. ⁷⁵
Plasmid: pCAGGS-PB-mCherry-CAXX	Gift from Dr. Cagla Eroglu	N/A
Plasmid: pCAGGS-EX	Gift from Dr. Nicholas Gaiano	Mizutani et al. ⁷⁴
Plasmid: pCAGGS-EGFP-nls	This paper	N/A
Plasmid: pCAGGS-EGFP-nls-Arhgap11a 3'UTR	This paper	N/A
Plasmid: pCAGGS-EGFP-nls-Arhgap11a 5'UTR	This paper	N/A
Plasmid: pCAGGS-EGFP-nls-Arhgap11a CDS	This paper	N/A

(Continued on next page)

Continued

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Plasmid: pCAGGS-Dendra2	Dr. Debra Silver Lab	Pilaz et al. ²¹
Plasmid: pCAGGS-Dendra2-Arhgap11a-noUTR	This paper	N/A
Plasmid: pCAGGS-Dendra2-Arhgap11a-5'UTR	This paper	N/A
Plasmid: pCAGGS-5'UTR-Arhgap11a-Dendra2	This paper	N/A
Plasmid: pCAGGS-Arhgap11a(GD)-Dendra-5'UTR	This paper	N/A
Plasmid: pCAGGS-CFP-MS2-noUTR	Dr. Debra Silver Lab	Pilaz et al. ²¹
Plasmid: pCAGGS-CFP-MS2-Arhgap11a 5'UTR	Dr. Debra Silver Lab	Pilaz et al. ²¹
Software and algorithms		
FIJI	NIH	https://ImageJ.NIH.gov/IJ
GraphPad Prism (9)	GraphPad	https://www.graphpad.com/scientific-software/prism/

RESOURCE AVAILABILITY**Lead contact**

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Debra Silver (debra.silver@duke.edu).

Materials availability

All unique/stable reagents generated in this study are available from the **Lead Contact** either without restriction or in some cases with a completed Materials Transfer Agreement.

Data and code availability

This study did not generate any unique code. Original/source data for figures in the paper are available from the corresponding author on request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS**Mice**

All experiments were performed in agreement with the guidelines from the Division of Laboratory Animal Resources from Duke University School of Medicine and approved by Duke IACUC. Plug dates were defined as embryonic day (E) 0.5 on the morning the plug was identified. All experiments were conducted in the C57BL/6J strain. The following mouse strains were used: Dlx-Cre (Tg(*Dlx5a-cre*)1Mek/J); Ai14-tdTomato (B6.Cg-Gt(ROSA)26Sor^{tm14(CAG-tdTomatoHze)/J}) both from Jax labs). *Arhgap11a*^{em1(IMPC)J} mice were generated by obtaining cryopreserved sperm from the KOMP consortium. These were used to produce germline knockout mice which were validated by qPCR and western analyses.

Human embryonic samples and *in situ* hybridization

The study using human fetal sample (9 wpc) was approved by three relevant Ethics Committees (Erasmé Hospital, Université Libre de Bruxelles, and Belgian National Fund for Scientific Research FRS/FNRS) on research involving human subjects. Written informed consent was given by the parents in each case. *In situ* hybridization (ISH) on human fetal cortical sections using digoxigenin-labeled RNA probes was performed as described previously.⁷¹ The probe specifically recognizing exon11 and 12 of human *ARHGAP11A*, not *ARHGAP11B*, was prepared (2117–2657 of human *ARHGAP11A* (NM_014783.5)). Imaging was performed using a Zeiss Axioplan 2 and the intensity and contrast were modified using Fiji/ImageJ software. We confirmed the specificity of the signal produced by the anti-sense pan-*ARHGAP11A* probe by comparing with the virtual absence of the signal by the sense probe.

METHOD DETAILS**Histology**

Mouse embryos were collected and dissected in cold phosphate buffer saline (PBS) and histology was performed as previously.²¹ Brain fixation was performed overnight by immersion in a 4% Paraformaldehyde 1X PBS solution. Following fixation, embryonic brains were washed twice in cold PBS for 20min. Cryoprotection was performed by overnight immersion in a 30% sucrose (w/v) PBS solution. Following cryoprotection, brain were transferred into OCT medium. Cryosections were generated using a cryostat and deposited on glass slides. The thickness of the sections varied depending on the purpose of the experiments (10–20µm for

characterization of ARHGAP11A localization by immunofluorescence and *in situ* hybridization, 50 μ m for characterization of basal process morphology with 3D reconstructions, 30 μ m for analysis of cortical layering at post-natal stages).

Immunofluorescence was performed as described previously.²¹ Briefly, slides were left to thaw at room temperature (RT) for 10min. Then, they were washed by immersion in PBS for 10min, followed by permeabilization with a 15-30min wash in 0.2-0.5% Triton-X (w/v) in PBS. Following this step, sections were washed once in PBS and blocking was performed with Mouse on Mouse (MOM, Vector Laboratories) blocking agent when using primary antibodies produced in the mouse, and 10% NGS for primary antibodies generated in any other species. Primary antibody incubation in PBS or MOM diluent was then performed overnight at 4C. Three 5-10-minute washes were then performed in PBS, followed by 30-60min RT incubation in a secondary antibody solution containing Hoechst. Before mounting was performed in Vectashield (Vector Laboratories) or Mowiol, three 5-10-minute washes were performed in PBS. The list of primary antibodies used in this study can be found in [STAR Methods](#). Cell counting was performed in FIJI (ImageJ), using the *Cell counter* plugin.

For binning analyses, X-Y coordinates were extracted from *Cell counter* data, and a script created in R⁷⁶ was used to assign punctae to specific bins, using the coordinates of the ventricular and pial borders as references. Bin numbers were reported in an Excel spreadsheet for analyses.

DNA constructs and siRNAs

pCAGGS-EX and pCAGGS-EGFP plasmids were kind gifts from Nicholas Gaiano,⁷⁴ the pGLAST-EGFP-CAXX plasmid was kindly offered by Tarik Haydar,⁷⁵ and the pCAGGS-PB-mCherry-CAXX was generously given by Cagla's Eroglu laboratory. EGFP-nls-CDS, EGFP-ns-5'UTR and EGFP-nls-3'UTR constructs were generated by cloning sequences of interest downstream of the EGFP sequence, using *Arhgap11a* cDNA cloned from mouse embryonic cortical cDNA as described previously.²² The MS2-no UTR plasmid was described previously.²¹ We used a Gibson assembly strategy (NEB Hifi Builder) to clone *Arhgap11a*'s 5'UTR from the EGFP-nls-5'UTR construct into the MS2-no UTR plasmid downstream of the MS2 stem loops sequence. Similarly, we used Gibson assembly to generate the Dendra2-no UTR and the Dendra2-5' UTR plasmids, cloning 2 fragments (*Arhgap11a* and *Dendra2* coding sequences) or three fragments (*Arhgap11a* and *Dendra2* coding sequences followed by *Arhgap11a* 5'UTR) into the EcoRI site of pCAGGS-EX, using EGFP-nls-CDS, Dendra2-no UTR,²¹ and EGFP-nls- 5'UTR as templates, respectively. siRNAs targeting the 3'UTR of *Arhgap11a* were purchased (siRNAflex, Qiagen), and a pool of 3 different siRNAs were used in these assays. These siRNAs were previously validated by the manufacturer.

In utero electroporation (IUE)

We performed IUE as described previously.^{21,77} Electroporation parameters were as follows: five consecutive 50ms electrical pulses spaced by 950ms, voltage varied from 40V to 60V depending on the embryonic stage at which the procedure was performed. Plasmids were produced using Qiagen or Sigma Endotoxin Free Maxi Prep kits and following the manufacturers' instructions. Individual plasmid concentrations injected into lateral ventricles ranged from 0.5 to 1 μ g/ μ l. siRNAs were injected at a final concentration of 2.5 μ M.

Single molecule fluorescent *in situ* hybridization (smFISH)

In situ hybridization was performed following the protocol described by Takahashi and Osumi.⁷⁸ Probes sequences are listed in [STAR Methods](#). The protocol used to reveal *Arhgap11a*, *MS2* or *Dendra2* mRNAs by smFISH was as previously.²¹ All buffers and solutions used for this protocol were previously treated with diethyl pyrocarbonate to quench RNA-ase activity. A pool of Stellaris probes targeting *Arhgap11a* and labeled with the Quasar570 fluorophore were purchased from Biosearch. *MS2* and *Dendra2* probes were described previously.²¹ For quantification of *Arhgap11a* mRNA density after IUE-mediated siRNA knockdown, 10 μ m mosaic Z-stacks covering the entire thickness of the electroporated and the corresponding region in the non-electroporated hemisphere were acquired using a 63X objective with a microscope equipped with Apotome technology (Zeiss). The electroporated region was evident based on the presence of EGFP+ cells. Coordinates of smFISH punctae were manually registered using the *Cell Counter* plugging in FIJI (ImageJ, over one thousand punctae were registered in non siRNA treated regions). A script created in R⁷⁶ was used to assign punctae to specific bins. Bin numbers were reported in an Excel spreadsheet.

qPCR analyses in *Dcx*-DsRed embryos

Cortices from E14.5 *Dcx*::DsRed embryos were isolated, incubated with 0.25% trypsin-EDTA solution for 10 min. at 37°C, dissociated into a single cell suspension, and sorted in a Sorter Astrios machine. Positive and negative cells were directly collected into RNA extraction buffer (RLT) supplemented with 1% β -Mercaptoethanol. Samples were vortexed and RNA was extracted using Qiagen RNeasy plus kit. cDNA was synthesized from RNA using Biogen iScript kit and qPCR was performed using either Sybr Green iTaq (BioRad) or TaqMan (Life Technologies) in an Applied Biosystems StepOne machine (Thermo Fisher Scientific). The following primers and TaqMan probes were used in the qPCR reaction: β -Actin (5' Forward-AGATCAAGATCATTGCTCCT and 3' Reverse-CCTGCTTGCTGATCCACATC), *Pax6* (5' Forward- TCTTTGCTTGGGAAATCCG and 3' Reverse-CTGCCGTTCAACATCCT TAG), *Arhgap11a* (5' Forward-GCAGGTGTGCCAAGGCGAAGT and 3' Reverse-TGCAAGTCGCCAACCAACACTTTCA)²⁸, *Gapdh* (Mm99999915_g1), *Tubb3* (Mm00727586_s1). Values were normalized to *Gapdh* (TaqMan) or β -Actin (Sybr Green) as loading control.

Live imaging

We performed and quantified live imaging of RNA trafficking, using methods identical to those described previously.²¹ For imaging of translation, following *in utero* electroporation of *Dendra2* and pCAGGS-CFP plasmids at E15 in the afternoon, brains were dissected in cold 1x HBSS supplemented with 2.5 mM HEPES, 30 mM D-glucose, and 4mM NaHCO₃ during the morning on the next day. We generated endfoot preparations as described previously,²¹ using tweezers to peel off the basement membrane together with connected endfeet from the surface of the brain. 2-4 endfoot preparations from brains electroporated with different plasmid conditions were mounted together in a 1mg/ml collagen solution supplemented with DMEM/F12 at the bottom of 35-mm glass bottom dishes (MatTek). A slab of 3% agarose gel was added on top of endfoot preparations to prevent their detachment from the bottom of the dish. Additional collagen was added to stabilize the preparation. Endfoot preparations were cultured in DMEM-F12 supplemented with B27 without vitamin A (Gibco), N2 (Gibco), 5% horse serum, 5% fetal bovine serum, 10ng/ml FGF and 20ng/ml EGF. Culture medium was added after a 15min incubation at 37C to ensure proper polymerization of collagen matrix. Endfoot preparations were left to equilibrate at 37C and 5% CO₂ for 1-2h prior to live imaging. Live imaging was performed with a 100x/1.4 oil U PlanSApo objective mounted on an inverted spinning disk confocal microscope (Andor XD revolution spinning disk confocal microscope), equipped with a 37C and 5% CO₂ incubation chamber. Following a one-hour equilibration of the incubation chamber, three 15 μ m Z-stacks per endfoot preparations were imaged in the blue, green and red channels with a 2 μ m resolution in the Z axis (pre-conversion acquisitions). This allowed the simultaneous recording of several conditions within one session and therefore minimized variability between imaging sessions. Positions were selected at locations where endfoot preparations presented minimal folding. Each position was then exposed to Arc lamp illumination at 10% intensity for 20 seconds with manual scanning in the Z-dimension. Each Z-stack was then acquired in the blue, green and red channels (post conversion acquisitions). Z stacks were then acquired every 5min for 45min in the green and blue channels (timecourse acquisitions). Following this, 40 μ M anisomycin treatment was performed for 20min and positions unaffected by the initial photoconversion were imaged using parameters described above (Aniso acquisitions). The FIJI software was used for the quantification of green fluorescence signal recovery over time. Z-projections were generated for all the Z-stacks. The ellipse-selection tool was used to generate regions of interests (ROIs) covering individual endfeet at each time point. Average green fluorescence intensity in endfeet was reported at each timepoint into an excel table where *Dendra2* recovery was calculated using the following formula: $(Dendra2\ recovery)_t = ((Dendra2\ signal)_t - (Dendra2\ signal)_{t_0}) / (Dendra2\ signal)_{t_0}$.

Analyses of basal process branching in the MZ

Slides containing sections from brains electroporated with pGLAST-EGFP-CAXX, siRNAs and rescue plasmids were washed once in PBS for 10min, followed by a 15min wash in 0.25% Triton X in PBS (w/v) and one additional PBS wash for 15min. Vectashield was used to mount the slides with a coverslip and prevent bleaching of the EGFP signal. 20-40 μ m Z-stack images were acquired at a 0.2 μ m resolution in the Z dimension, using a 63X objective mounted on an epifluorescence microscope equipped with the Apotome technology (Zeiss). Analyses were performed in FIJI. Maximum intensity Z-projections of the Z stacks were generated to identify entire basal processes and endfeet. Rectangular regions of interests (ROIs) were traced around individual or small groups of basal processes. 3D projections of selected ROIs were generated with the following parameters deviating from the default settings: "axis of rotation": Y axis, "rotation angle increment": 5, "Interpolate": checked. We used the *Cell Counter* plugin to count branches of each order, rotating the 3D projection in order to identify branches obscured by other basal processes. First order branches were those located further away from the pial surface (see Figure 4 for a depiction of branch orders). We used the *Line* tool to ensure that we counted branches >5 μ m only. The number of branches of each order was then reported into an Excel spreadsheet and the total number of branches was calculated as the sum of all the branches from different orders.

Analyses of endfoot area covering the basement membrane

These analyses were performed in fixed endfoot preparations from electroporated brains. These preparations were mounted in mounting medium on a glass slide, covered with a coverslip. We were careful to ensure that the EGFP+ endfeet were located immediately under the coverslip. Native EGFP fluorescence was acquired in Z-stacks with a confocal microscope. Using Maximum Intensity Z-projections, we employed the *Polygon Selection* tool in ImageJ to trace the outside limits of basal endfeet, and measured areas were reported into an Excel spreadsheet.

Analyses of basal process extensions in the IZ-low CP regions

These analyses were very similar to those performed in the upper CP/MZ regions, with the following modifications. Z-stack images of basal process in the IZ, low CP regions were acquired on a LSM 710 Zeiss confocal microscope. Here we did not assess the order of branches and a 5 μ m cutoff was not used. Instead we counted the total number of extensions emanating from the acquired region of the basal process and normalized this number to the length of basal process that we acquired and focusing on processes with a visualized length >50 μ m. Rotation of 3D projections were still used to visualize extensions masked by other structures.

Live imaging of neuron migration

A first *in utero* electroporation (IUE) of a pCAGGS-EGFP plasmid was performed at E14.5. 24h later, another IUE procedure was performed to transfect siRNAs together with a pCAGGS-PB-mCherryCAAX into the hemispheres transfected earlier with

pCAGGS-EGFP. In the morning of the next day, 300 μ m-thick organotypic brain slices were generated as described previously.⁷⁹ Slices were cultured on cell culture inserts, and were surrounded with a 1mg/ml collagen solution supplemented with DMEM-F12. We used the same medium as the one described above for Dendra2 live imaging of basal endfeet. 4-6 slices transfected with different siRNA conditions were mounted on the same inserts. In the evening, live imaging was initiated for a total of 16h. Z-stack images of regions showing clear overlap between the first and second electroporations were acquired every 10min, using a 10x objective mounted on an inverted LSM 710 Zeiss confocal microscope, equipped with an incubation chamber (37C, 5% CO₂). Analyses were performed in FIJI, using the *Manual Tracking* plugin. We focused on neurons located in the low IZ / SVZ region at t_{0h} . Tracking results were reported in an Excel spreadsheet and the analyzed parameters are defined as follows. Net distances in Y and X: $|Y_{0h}-Y_{16h}|$ and $|X_{0h}-X_{16h}|$, respectively. Net distance: Euclidean distance between neuron positions at t_{0h} and t_{16h} . Total path length: sum of distances traveled between each time points. Velocity: average travel speed between each time point. For analysis of movements in Y as “down”, “steady”, or “up”, we calculated the Y distance traveled from one time point to the next ($Y_{t1}-Y_{t2}$). “Down”, “steady” and “up” were defined as negative, null and positive Y distances, respectively.

Serial block face electron microscopy

For SBF-SEM, the samples then underwent a heavy metal staining protocol adapted from Deerinck (2010) after fixation. The tissues were washed in 0.1M sodium cacodylate pH7.4 and then a solution of 1.6 % potassium ferrocyanide with 2% osmium tetroxide buffered with 0.1M sodium cacodylate was added for 1h at RT. This is followed by filtered 10% thiocarbonylhydrazide (TCH) freshly prepared for 30min. Samples were then washed in distilled water and a secondary 2% osmium tetroxide incubation for 30min. The samples were then placed in 1% uranyl acetate at 4C overnight, washed in distilled water and then placed in freshly prepared lead aspartate solution for 30min at 60C. The samples were then dehydrated with cold ethanol, from 25% to 100% and then infiltrated with increasing concentrations of Durcupan resin in ethanol with several exchanges of 100% resin. The samples were finally embedded in 100% resin and left to polymerize at 60°C for 48 hours.

The tissue samples embedded in resin were manually trimmed with a razor blade to expose the tissue on their surfaces, and then glued onto an aluminum SBF-SEM rivet with conductive epoxy (SPI Conductive Silver Epoxy) with the exposed tissue down. Specimens on the rivet were further trimmed with a razor blade to as small a size as possible (about 0.5mm), and block face was trimmed with a glass knife. Once tissue was exposed, semi thin sections 0.5 μ m were taken and stained with toluidine blue and viewed under a light microscope to check tissue orientation and condition. Then, the rivet with the sample was sputter coated with gold-palladium.

The image stacks were acquired in an automated fashion by using a high resolution scanning electron microscope (Merlin - Carl Zeiss, Germany) equipped with a 3View system (Gatan Inc., Pleasanton, CA, USA), and a back-scattered electron detector. The Digital Micrograph software (Gatan Inc.) was used to adjust the SEM imaging conditions and slicing parameters. The electron microscope was operated at the high-resolution mode with an acceleration voltage of 2 kV, current mode and in the high-vacuum mode. All images were taken with the following scanning settings at 80pA, dwell time = 2s; pixel sizes 5-7nm. On average 300 sections were obtained at 60nm thickness

QUANTIFICATION AND STATISTICAL ANALYSIS

All analyses were performed in a blinded fashion. Number of data points and statistical tests used for all the comparisons are indicated in the figure legends.