

HHS Public Access

Wiley Interdiscip Rev RNA. Author manuscript; available in PMC 2015 October 28.

Published in final edited form as:

Author manuscript

Wiley Interdiscip Rev RNA. 2015 ; 6(5): 501-515. doi:10.1002/wrna.1289.

Post-transcriptional regulation in corticogenesis: how RNAbinding proteins help build the brain

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Abstract

The cerebral cortex, the brain structure responsible for our higher cognitive functions, is built during embryonic development in a process called corticogenesis. During corticogenesis, neural stem cells generate distinct populations of progenitors and excitatory neurons. These new neurons migrate radially in the cortex, eventually forming neuronal layers and establishing synaptic connections with other neurons both within and outside the cortex. Perturbations to corticogenesis can result in severe neurodevelopmental disorders, thus emphasizing the need to better understand molecular regulation of brain development. Recent studies in both model organisms and humans have collectively highlighted roles for post-transcriptional regulation in virtually all steps of corticogenesis. Genomic approaches have revealed global RNA changes associated with spatial and temporal regulation of cortical development. Additionally, genetic studies have uncovered RNA-binding proteins (RBPs) critical for cell proliferation, differentiation, and migration within the developing neocortex. Many of these same RBPs play causal roles in neurodevelopmental pathologies. In the developing neocortex, RBPs influence diverse steps of mRNA metabolism, including splicing, stability, translation, and localization. With the advent of new technologies, researchers have begun to uncover key transcripts regulated by these RBPs. Given the complexity of the developing mammalian cortex, a major challenge for the future will be to understand how dynamic RNA regulation occurs within heterogeneous cell populations, across space and time. In sum, post-transcriptional regulation has emerged as a critical mechanism for driving corticogenesis and exciting direction of future research.

INTRODUCTION

The cerebral cortex is the most complex biological "machine" known to man. Part of this complexity resides in the web of coordinated functional units, the cortical areas. Cortical areas are radially organized within layers, each of which contain neurons with similar

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Conflict of interest: The authors have declared no conflicts of interest for this article.

molecular, electrophysiological, and connectivity characteristics.¹ The cytoarchitecture of an area and thus the number of neurons in each layer is paramount to specify its post-natal function. Additionally, glial cells (astrocytes, oligodendrocytes, and microglia) play a key role in the homeostasis of the cortex. Defects in cortical development can cause acute neurological disorders affecting brain size and function such as microcephaly or lissencephaly. Therefore, the developmental mechanisms that regulate neuronal number and positioning together with glial cells are crucial to build a healthy brain. This review will focus on the molecular regulation of neuronal generation and positioning during embryonic neocortical development.

During embryonic development, excitatory neurons are generated from neural progenitor populations in a process termed neurogenesis.² The germinal zones include the ventricular zone (VZ), located at the border of the cerebral ventricles, and the subventricular zone (SVZ) located beside the VZ (Figure 1). During early cortical development the predominant neural progenitors are neuroepithelial cells (NE cells), which mainly undergo symmetric proliferative divisions to self-renew. NE cells are later replaced by radial glial cells (RGCs), which primarily undergo asymmetric divisions to generate a new RGC and a more differentiated cell, either a neuron or an intermediate progenitor (IP)^{3,4}. Both NEs and RGCs extend processes from the ventricular border to the pial surface while their cell body (nucleus) resides in the VZ. IPs are lineage-restricted multipolar progenitors which divide in the SVZ to amplify the neuronal population. $^{5-7}$ Hence neurons are directly generated by both RGCs and IPs. In mice, the most widely utilized model for studying corticogenesis, the proliferative period begins around embryonic day (E) 10.0, and the neurogenic period begins about E11.5 and continues to E18.5. Neurons of different layers are born in a sequential fashion, with deep layer neurons born between E11.5 and E14.5 and superficial layer neurons born between E14.5 and E18.5.8,9

After their generation, newborn neurons migrate toward the pial surface of the cortex, using the basal process of RGCs as their scaffold. Their route passes across the intermediate zone (IZ) in the middle of the cortex and ends in the cortical plate (CP), the final location of neuronal layers (Figure 1). During normal development young neurons migrate up to the pial surface, bypassing neurons born earlier.^{10,11} Thus, deep layer neurons born earlier in development eventually reside closer to the ventricle, whereas superficial layer neurons are ultimately found near the pial surface. Upon reaching their final position within the cortex, the excitatory neurons then establish connections with other neurons both within and outside of the cortex. Hence the fate and final function of projection neurons is ultimately defined by their birth and subsequent migration to distinct layers of the brain.

Although much is known about transcription factors and signaling molecules in corticogenesis, only recently have we begun to appreciate the widespread roles of RNAbinding proteins (RBPs) in neocortical development. Specific RBPs are expressed across different developmental stages of the developing neocortex as evidenced by quantitative RT-PCR analyses, *in situ* hybridization and transcriptome – analyses.^{34–36} However only a small number of RBPs have actually been tested for a functional impact upon corticogenesis (Table 1). Those RBPs important for neocortical development impact diverse steps of RNA metabolism, and thus collectively reveal that modulation of all stages of the RNA life cycle

is necessary for cortical development (Figure 2). In this review we highlight critical RBPs implicated in embryonic corticogenesis, including the production, differentiation and migration of excitatory neurons. We describe their known functions in RNA regulation, cortical development, and in relevant cases disease pathogenesis. These include both RBPs strictly expressed in the developing cortex and ubiquitous RBPs with enriched neocortical expression. We organize our review by discussing neocortical RBPs relevant for each major step of posttranscriptional regulation: alternative splicing, nucleo-cytoplasmic transport, RNA stability and translation, and localization (Figures 2 and 3).

ALTERNATIVE SPLICING

Alternative splicing (AS) is a powerful mechanism to amplify the output diversity of the genome through the editing of primary transcripts (Figure 2). The excision or inclusion of intronic and exonic sequences of pre-mRNA produces distinct transcripts that may be translated into biochemically diverse proteins. AS of the 5' and 3' UTR, or coding regions of a pre-mRNA can also impact downstream steps of mRNA metabolism including stability, nuclear export, nonsense mediated decay (NMD), and RNA localization, largely by exposing binding sequences for RBPs or miRNAs. Hence, AS is a powerful mechanism to differentially manipulate gene expression between cells, and recent studies reviewed below underscore its relevance for cortical development.

Genomic Analyses of AS in the Developing Neocortex

With the advent of transcriptome techniques, we are now beginning to appreciate the contribution of genome-wide RNA splicing for cortical development. Studies of both mouse and human cortical models have collectively revealed both spatial and temporal AS differences during cortical development (Table 2). One of the first studies to dissect AS came from Nenad Sestan's group, who utilized whole-genome exon microarrays to reveal region specific differences in AS in the human brain at mid-gestation.³⁷ This comprehensive study discovered that at mid-fetal development, 28% of expressed genes are alternatively spliced between different human brain structures. Among those genes showing robust AS is *ROBO1*, which is involved in axon guidance and neural progenitor proliferation, and is implicated in various neurodevelopmental disorders.^{38,39} Distinct AS *ROBO1* transcripts might promote establishment of connectivity and/or the generation of the appropriate number of neurons in distinct layers of mature cortical areas.

A similar spatial analysis of AS was applied within the developing embryonic mouse cortex at mid-gestation (E14.5) by Ayoub et al., who coupled RNA sequencing with laser-capture microdissection to demonstrate the existence of differential AS between different embryonic cortical zones (VZ vs SVZ + IZ vs CP).³⁶ This study revealed that some genes, such as *Wdr61*, show no significant difference in overall expression levels between zones but do exhibit differential expression of splice variants. For other genes, such as *Mfge8*, differential expression of just one splice variant across cortical zones can explain overall shifts in expression. In addition for other classes of genes, such as *Cugbp2* and *Hes6*, the relative ratio of spliceoform expression is similar between zones indicating AS may be less relevant. The three embryonic cortical zones assessed by Ayoub et al. contain largely different

populations of progenitors and post-mitotic neurons. These spliceoform expression differences imply there are cell-specific AS within the developing mouse neocortex.

As embryonic development proceeds, the repertoire of progenitors and neurons also changes. Hence it is not surprising that in addition to spatial differences in AS, temporal differences in AS are evident across different stages of cortical development. Dillman et al. compared cortical samples from embryonic day (e) 16 to those from postnatal day 30 in the mouse, and discovered AS differences. Amongst these they noted that spliceforms more highly expressed postnatally encode actin-related proteins.⁴¹ This finding is of interest as actin metabolism is paramount in maturing neurons during early postnatal stages when neuronal connections are being established.⁴³ Temporal patterns of AS have also been discovered using *in vitro* models of corticogenesis, in which human embryonic stem cells are differentiated into neurons.⁴² This longitudinal analysis (termed Cortecon by the authors) revealed widespread AS genes were associated with cancer or nervous system diseases.

A significant limitation of the aforementioned analyses is that most samples analyzed to date contain heterogeneous cell populations, which collectively may contribute to AS differences, thus complicating interpretation. A recent comprehensive study by Molyneaux et al. significantly overcame this hurdle, using deep sequencing to probe transcriptome and AS changes in sorted excitatory neuronal populations from various stages of corticogenesis.⁴⁰ This group discovered 1181 genes with shifts in isoform expression during corticogenesis. From transcriptome analyses the authors identify genes showing uniform expression at the gene-level but significant differences at the isoform level, an observation also made by Ayoub et al. as described above for *Wdr61*.³⁶ Altogether, these analyses provide evidence that AS is at play in the developing cerebral cortex across multiple dimensions (tangential, radial, and temporal). The current studies collectively highlight new candidate genes that may regulate corticogenesis. Future studies that similarly apply cell sorting and/or single cell transcriptome analysis will be valuable for further discovery of AS differences in cortical development.

SPLICING FACTORS REGULATING CORTICOGENESIS

The spatio-temporal regulation of AS relies on the differential expression and function of trans-splicing factors including RBPs. Both McKee et al. and DeBoer et al. uncovered splicing factors expressed within the developing cortex.^{34,35} Several of these RBPs have been experimentally shown to be critical for cortical development.

NOVA2

NOVA1 and 2 are members of KH-domain RBPs and are among the best-characterized RBPs in the brain. Nova proteins bind RNA via YCAY clusters and UCAU sequences, and regulate AS *in vitro*. Of these, *Nova2* is highly expressed in the neocortex.⁴⁴ Consistent with biochemistry data, *Nova2* knock out mice contain a significant number of splicing anomalies in the postnatal brain.¹² Interestingly, the vast majority of mis-spliced genes (34 of 40) encode proteins localized to the synapse. During prenatal cortical development, NOVA2

binds to a large number of transcripts (27,576).¹³ Yano et al. showed *Nova2* is necessary for the proper migration of upper-layer neurons toward the CP. The authors used HITS-CLIP of *Nova2*-deficient brains to identify key downstream splicing targets of Nova2. They focused on 20 genes of the Reelin pathway, because Reelin is a migration cue secreted by early born Cajal Retzius (CR) neurons. Remarkably, the authors identified significant splicing changes in only one transcript of this pathway, finding that NOVA2 regulates excision of exons 7b and 7c of *Dab1*. Using *in utero* electroporation of minigene constructs, the authors then elegantly showed that *Nova2* is essential for the proper expression of *Dab1* spliceforms and that this splicing mediates neuronal migration in the neocortex. In the future it will be of interest to identify additional splicing targets genome-wide that may also be regulated by *Nova2* in the developing cerebral cortex.

PTBP2

Poly-pyrimidine tract-binding proteins (PTBP proteins) are involved in multiple steps of RNA metabolism including splicing. Biochemical studies of PTBP1 demonstrated this family of proteins binds introns (recognizing CU-repeats and UCUY-rich elements).⁴⁴ Depending on its relative positioning with respect to certain exons (either upstream or downstream), PTBP1 can either promote or inhibit exon inclusion.¹⁶ Although *PTBP1* is minimally expressed in the brain, PTBP2 (also called nPTB) is highly expressed in the brain.¹⁶ In the developing mouse brain, PTBP2 binds thousands of RNAs.¹⁶ Among these mRNAs, splicing of *Psd-95*, which encodes a synaptic protein, is repressed by PTBP1 and PTBP2 during development.¹⁷ To further understand the role of PTBP2 in cortical development, two groups recently generated PTBP2-deficient mice. Licatolosi et al. discovered that E14.5 PTBP2 mutant brains had defective neural progenitor polarity, accompanied by defects in proliferation and neuronal differentiation.¹⁶ Li et al. observed only postnatal cortical defects in their mutant mice, related to a role in neuronal differentiation, maturation, and survival.¹⁸ The phenotypic differences may be because of the nature of the mouse mutation, as the former study used germline knockouts and the latter study used conditional expression with Nestin-Cre and Emx1-Cre lines. Regardless, in both studies, PTBP2-deficiency was associated with significant alterations in AS, including mRNAs involved in regulation of the actin cytoskeleton, proliferation and cell fate. Although RNA-targets of PTBP2 involved in the regulation of neurogenesis have not yet been identified, these studies establish PTBP2 as a key regulator of AS in both neural progenitors and immature neurons.

TRA2B

TRA2B splicing factor is a member of the Serine-/Arginine-Rich (SR) protein family, which have well-established roles in constitutive and AS.⁴⁵ SR proteins recognize exonic splice enhancers, as well as interact with other splicing factors to promote splice site recognition. Once splicing is complete, SR proteins may or may not remain on the mRNA. This retention of SR proteins on the mature mRNA can impact nuclear export and downstream RNA regulation. TRA2B regulates splicing of *Tau* and *Smn2* mRNAs, involved in Alzheimer's disease and spinal muscular atrophy, respectively.^{46,47} Hence *PTBP2* is relevant for human diseases of the nervous system. Constitutional loss of *Tra2b* in the mouse leads to

embryonic lethality by E7.0, precluding the use of this model for the study of its role in cortical development.⁴⁸ However conditional knock-out embryos using either *Emx1*-Cre¹⁴ or *Nestin*-Cre¹⁵ point toward an essential role of *Tra2b* in the survival of neural progenitors and neurons. Strikingly, conditional *Tra2b* knockout in the cerebral cortex leads to almost complete absence of the cortex at adulthood, following a massive wave of apoptosis during embryonic cortical development. These studies establish the fundamental requirement of *Tra2b* in survival of neural cells, and highlight the importance of future studies to determine key mRNA targets of TRA2B in the embryonic cortex. Altogether, these data suggest that AS regulation plays a critical role in cortical development. Given the abundance of splicing factors in the developing brain, clearly these studies are just the tip of the iceberg.

From the Nucleus to the Cytoplasm: The Exon Junction Complexes

As splicing proceeds, spliced transcripts become decorated by exon junction complexes (EJC), which bind primarily at the junctions where introns are excised.^{49,50} The EJC remains bound to the spliced mRNA as the RNA is exported into the cytoplasm. The core EJC is composed of the heterodimer, Magoh and Rbm8a, the helicase, Eif4a3, and the cytoplasmic component Casc3. This core complex interacts transiently with other proteins to mediate various aspects of mRNA metabolism, including mRNA splicing, localization, nonsense-mediated decay, and – translation initiation.^{51–53}

Magoh was recently shown to be essential for corticogenesis in mice.^{19,54} Haploinsufficiency for *Magoh* causes a severe microcephaly, associated with depletion of IPs and massive apoptosis of new neurons. Silver et al. showed that Magoh regulates proper cell division of radial glia and hypothesized that defective mitosis induces aberrant production of progenitors and neurons.¹⁹ Microarray analysis of *Magoh* haploinsufficient cortices revealed only 147 transcripts with significant differential expression. Given that the EJC decorates >80% of exon–exon junctions⁴⁹ this indicates *Magoh* haploinsufficiency may not globally impact RNA stability although this remains to be formally tested. The authors identified protein changes downstream of Magoh, including one physiologically relevant target, Lis1, a microtubule-associated protein also involved in brain development. Future studies will be useful to assess how Magoh impacts radial glia divisions either via translation and/or some other step in mRNA metabolism.

The same group recently probed whether haploinsufficiency for the Magoh heterodimer, Rbm8a, impacts corticogenesis.²⁰ Similar to *Magoh, Rbm8a* is also highly expressed in the developing cortex. Conditional haploinsufficiency for *Rbm8a* induced microcephaly, even more severe than *Magoh* loss. This phenotype was associated with depletion of progenitors and dramatic apoptosis especially of neurons. *Rbm8a* mutant embryos showed precocious neuron production and faster cell cycle exit of progenitors. Thee phenotypic similarities induced by *Magoh* and *Rbm8a* haploinsufficiency support a model where Magoh and Rbm8a act together as part of the EJC to regulate corticogenesis. The observed differences in severity of phenotypes could reflect distinct functions outside of the EJC or redundancies with other proteins, such as *MagohB*.⁵⁵ In addition to these roles of core EJC components in the brain, the peripheral EJC component involved in NMD, Upf1, is also expressed in the developing neocortex and promotes a stem cell state in primary cells.⁵⁶ Future genetic and

molecular studies of these mutants will help establish which aspect(s) of EJC regulation are critical to development of the brain.

Recent mouse and human genetic studies have collectively implicated EJC dosage in neurodevelopmental pathologies associated with aberrant cortical development, including autism, schizophrenia, and – intellectual disability.^{57–59} UPF3B, an EJC component required for nonsense-mediated decay, is mutated in X-linked intellectual disability, schizophrenia, and autism. Copy number variations in several EJC components, including *UPF3B, EIF4A3, RBM8A*, and *MAGOH*, are found in patients with intellectual disability frequently accompanied by brain malformations.⁵⁷ *RBM8A* is within the proximal 1q21.1 microdeletion/duplication associated with microcephaly.⁵⁸ *EIF4A3* is mutated in Richeiri-Costa Syndrome, a developmental disorder which can also be associated with brain malformations.⁶⁰ Continuing genetic studies in model organisms will help establish if roles for these EJC components in corticogenesis are the root causes for these neurodevelopmental disorders.

RNA Stability and Translational Control

Outside of the nucleus, RNA stability and translational regulation offer yet another layer of control for gene expression. The role for RNA stability in corticogenesis is poorly defined. A number of RNAs have been shown to have short-half lives but so far this has been attributed to oscillations in transcription. Translational control can be exerted at different steps: initiation, elongation, or termination. These steps are regulated by the coordination between ribosomal complexes and a vast set of RBPs. RBPs which impact translation control have recently been shown to influence development of the cerebral cortex.

HuR

HuR is a well-characterized RBP, part of a family of Hu-related proteins that preferentially bind the 3'UTR of its RNA targets to influence multiple aspects of RNA metabolism including RNA stability and translation.^{61,62} While several Hu proteins have been implicated in neuron differentiation and post-mitotic function,^{63,64} so far only HuR has been formally shown to regulate normal embryonic corticogenesis. *HuR* is expressed early in neuroepithelial cells, when these progenitors are undergoing primarily proliferative divisions.²¹ Garcia-Dominguez et al. postulated HuR influences the Notch pathway by regulating mRNA levels of the ligand Delta.²¹ The expression of Delta ligand promotes proliferation and prevents differentiation in neighboring cells via a mechanism called lateral-inhibition.⁶⁵ The authors discovered that HuR interacts with *Dll1* mRNA. *HuR* depletion in neural precursors leads to reduced *Dll* mRNA levels and less differentiation. This differentiation phenotype can be rescued by overexpression of *Dll1*. Hence the authors propose that HuR regulates *Dll1* stability to promote lateral inhibition and thus influence early cell fates in the cortex.

Later in development when neuroepithelial cells have been replaced by radial glial progenitors, HuR is expressed in radial glia, IPs, and newborn neurons.²² In an elegant study evaluating translation, Kraushar et al. used genome-wide polysome-profile analysis in conditional *HuR* knockout murine cortices at E13 and P0 to uncover a large pool of *HuR*-

regulated mRNAs redistributed to different polysomal fractions during development. HuRdependent RNAs were enriched for regulators of transcription, translation and layer specific pathways. The transcripts regulated by HuR were dramatically different in E13 and P0 brains, perhaps reflecting different biological processes occurring at these distinct ages. The authors also make the novel discovery that HuR interacts with the Eif2 kinase, Eif2ak4, which regulates the presence of distinct ribosomal proteins in active sites of translation at polysomes. The authors argue that HuR coordinates the translation of a network of mRNAs encoding proteins that share common functions, akin to the RNA regulon model first proposed by Jack Keene.⁶² Consistent with a functional requirement of *HuR* in cortical development, phenotypic anatomical analyses of P0 *HuR* conditional knockout mice revealed that HuR regulates the position, identity and maturation of post-mitotic glutamatergic neurons. Future work will be valuable to further identify the molecular mechanisms by which *HuR* regulates these developmental processes. Moreover this study sets the stage for future identification of signals that influence temporal control of mRNA translation.

Musashi

The translational regulators, Musashi 1 and 2 (Msi-1 and Msi-2), are highly expressed in NSCs throughout the central nervous system, including the mammalian cortex,⁶⁶ Cortical $Msi-1^{-/-}$ dissociated NSCs transfected with antisense peptide-nucleic acid against Msi-2showed decreased neurosphere formation and proliferative capacity, perhaps linked to impaired cell-cycle progression.²³ Hence Msi-1 and Msi-2 have redundant functions in neural stem cells. Musashis (Msis) are reported to act as both positive and negative regulators of translation, effects that are mediated through binding to the 3'UTRs of target mRNAs, including mammalian Numb.²⁴ Although this translational relationship between Msi and Numb has so far been shown in fibroblasts it is tempting to speculate that it may also hold true in NSCs, where numb is important for influencing neurogenesis.^{67,68} Msi targets have not yet been identified in NSCs; however, a recent study discovered translational targets from primary NSCs overexpressing Msi, using Ribosome profiling and RNA-seq.²⁵ Among several transcripts with reduced translation efficiency was Jag1, a ligand for Notch receptors, as well as a number of RBPs including Prpf3, Kirrel3, Rbm22, and *Dhx37*. Interestingly these targets have abundant Msi-binding sites in their 3' UTRs, thus Msi is thought to directly bind these targets. Katz et al. also demonstrated that Msi overexpression impacted AS while not perturbing overall RNA levels extensively.²⁵ Because Msi is primarily cytoplasmic, these changes are thought to be a secondary consequence of translational regulation of splicing factors, and not because of a direct role in AS or RNA stability per se. With identification of Msi translational targets, it will be of interest in upcoming studies to assess the role of these Msi targets upon NSC behavior in the cortex.

FMRP

FMRP (Fragile-X mental retardation protein) is an RBP encoded by the *Fmr1* gene. FMRP has been largely characterized as a translational inhibitor. *Fmr1* null mutations result in the Fragile-X syndrome (FXS) in humans, which is the most prevalent intellectual disorder

caused by mutations in one single gene.^{69,70} Postnatally, FMRP localizes at the synapses between neurons, where it inhibits the translation of a subset of localized mRNAs encoding proteins involved in synaptic plasticity.^{69,71} In response to neuronal activity, FMRP translational inhibition can be alleviated to allow for local, fast and massive, production of proteins necessary for structural modifications of the postsynaptic compartment.^{69,72,73}

Recent studies of humans and mice support a role for *FMRP* in regulation of prenatal cortical development. An analysis of brain region volumes of 1-to 3-year-old boys with FXS showed that several cortical areas display enlarged gray matter volume, suggesting a possible regional increase in neurons.⁷⁴ Indeed, analysis of P5 *Fmr1^{-/-}* mouse pups revealed increased neuronal density in the somatosensory cortical area.²⁶ Moreover cultured NSCs derived from either pre- or postnatal *Fmr1^{-/-}* mice generate more neurons than those derived from comparably aged WT mice.²⁷ Interestingly, the density of Tbr2+ IPs is higher in *Fmr1-KO* cortices, compared to controls, indicating increased neurons could be produced by more IPs.²⁶ Hence FMRP may control neuron production either by regulating IP differentiation or IP generation from radial glial cells. In support of the latter possibility, Saffary et al. used *in utero* electroporation to knockdown *Fmr1* in the developing cortex and demonstrate FMRP is required for IP generation.²⁸ The authors identified the candidate mRNA-target *Profilin1 (Pfn1)* as a mediator of this process, finding that *Pfn1* over-expression rescues the overproduction of IPs in *Fmr1* mutant brains.

Another recent study revealed a role for FRMP in neuronal migration in the cortex. In a study by La Fata et al., newborn neurons labeled by *in utero* electro-poration in *Fmr1* knockout brains showed defective neuronal migration.²⁹ These defects eventually lead to abnormal neuronal networks in the postnatal brain, which could be rescued by the overexpression of N-Cadherin, an mRNA target of FMRP. In addition to its well-established role in translational repression, FMRP has also been implicated as a pro-translation regulator in young neurons of the human neocortex.³⁰ Kwan et al. showed that FMRP expression enhances the translation of *NOS1*, an important regulator of synapse formation and spine maintenance.^{30,75} Interestingly FMRP-mediated regulation of *NOS1* translation was not evident in mouse projection neurons, highlighting potentially interesting evolutionary differences in FMRP function. Altogether these two studies suggest that defective neuronal circuits induced by defects in immature neurons could be at the origin of Fragile-X pathology in the adult.

Pfn1, NOS1, and N-Cadherin are likely part of a vast FMRP-regulated mRNA network involved in the regulation of the NSC-to-IP transition, early neuronal differentiation, and migration. Although FMRP targets have been elucidated in adult brains,⁷⁶ it still remains an outstanding question which RNAs are FMRP targets in neural stem cells of the developing neocortex and whether their translation is repressed or activated by FMRP. Future work is needed to identify those potential targets, and to assess their contribution to behavior of NSCs and neurons.

Eif4E/4E-T Complex

The *EIF4E* protein family, composed of *Eif4E1*, 2, and 3 (Eukaryotic Initiation Factor 4E), is part of a supercomplex docked to the 5' cap of mRNAs.⁷⁷ Once bound to mRNAs this

complex can either promote or inhibit translation, depending on its composition. These functions are mediated via interactions with additional translation factors. For example, *EIF4E1* association with *EIF4G* initiates translation whereas *EIF4E1* binding to *4E-T* blocks translation or promotes mRNA decay by targeting mRNAs to P bodies.⁷⁸

A role for translational regulators in corticogenesis was recently revealed using in utero knockdown in embryonic brains.³¹ Yang et al. discovered that decreased levels of either *Eif4e1* or 4*E*-*T* in neural progenitors lead to more neurons and fewer neural progenitors. This is accompanied by an increase in the number of cells with high protein levels for Ngn1, Ngn2, and Neurod1, basic-Helix-Loop-Helix (bHLH) pro-neuronal transcription factors. The authors discovered that in neural progenitors, EIF4E1 binds to Neurog1 and Neurog2 and NeuroD1 mRNAs. These biochemical results, along with rescue experiments using constructs deficient in RNA binding or protein-protein interactions, collectively revealed the Eif4e1/4E-T complex may repress translation of key neurogenic transcripts. Conversely, knockdown of Eif4G in neural progenitors (a positive regulator of translation when associated with *EIF4E1*) promotes fewer neurons and more progenitors. Altogether, these results suggest that certain neural progenitors are predisposed to the generation of neurons through the transcription of proneural bHLH transcription factors, but are stalled in a proliferating state by EIF4E1/4E-T-mediated translational repression of these target mRNAs. The authors speculate that NSCs are preloaded with mRNAs encoding prodifferentiation factors; however, translation of these mRNAs is repressed by Eif4E1 binding. This is a compelling hypothesis and it will be exciting in the future to test this model. It will also be valuable to demonstrate the direct role of Eif4E1/4E-T on translation of key mRNAs in NSCs, to rule out potential roles in nuclear export or sequestration of mRNAs in RNA-processing bodies.

RNA Localization

RNA localization plays a critical role in neurons both pre- and postnatally. When paired with translational regulation, RNA localization allows for local protein synthesis in the cytoplasm in response to intra- or extracellular signals (see Buxbaum et al., for a recent comprehensive review on this topic).⁷⁹ In immature neurons, RNA localization and translation, in response to extracellular guidance molecules, is paramount for axon guidance and synaptic function.⁸⁰ In mature neurons, RNA localization at the synapse may be involved in the precise, fast response of cells to integrate signals from other neurons, in order to consolidate or suppress memories.^{72,81} Recent studies have now highlighted roles for mRNA localization in mammalian neural progenitors, within both the cell body and basal process.

RNA Localization to the Basal Endfeet

Recently, it was shown that mRNAs encoding *CyclinD2* accumulate in structures called basal endfeet, located at the end of basal process.⁸² CyclinD2 is an outstanding candidate for the maintenance of neural progenitor proliferation as it is a well-characterized G1-phase regulator, and G1 phase is strongly linked to neural progenitor proliferation (Figure 1).^{83,84} Osumi's group identified a region in the 3'UTR of the *Cyclin D2* mRNA that is sufficient for its translocation to the basal endfeet. They also used a GFP reporter construct to argue

that these mRNAs are locally translated, although diffusion of GFP proteins from the cell body could not be ruled out from this experiment. During asymmetric division of NSCs, the daughter cell which adopts proliferative behavior inherits the basal process, whereas the daughter cell that does not inherit the basal process proceeds toward differentiation.⁸⁵ This led to the hypothesis that following cell division, *Cyclin D2* mRNA is locally translated and newly generated proteins subsequently migrate back to the soma through the basal process to promote proliferation. Identification of the transmachinery, including RBPs that bind *CyclinD2* will be useful for understanding why it is asymmetrically localized in NSCs, and for identifying additional localized RNAs.

Stau2

Stau2 is a double-stranded RBP, which in neurons has been well characterized as a translational repressor and a regulator of subcellular localization.⁸⁶ In neurons, Stau2positive RNA granules aggregate to form heterogeneous RNA granules that subsequently associate with motor proteins to translocate along microtubules to distal regions. Inspired by the Drosophila literature which established a role for Stau in neuroblasts, two independent groups recently showed that Stau2 plays a key role in mammalian cortical development.^{32,33} In mitotic progenitors, Stau2 is enriched at one pole of the cell, and becomes asymmetrically localized to only one of the post-mitotic progeny. Kusek et al. showed Stau2 was specifically inherited by the daughter cell that differentiates into an intermediate precursor cell following mitosis.³³ Both groups showed that downregulation of *Stau2* by shRNA knockdown leads to increased differentiation and a depletion of radial glia both in vitro and in vivo. Additionally, Vessey et al. showed that Stau2 acts in coordination with at least two other RBPs: the helicase Ddx1 and the translational repressor Pum2.³² RIPs were employed to identify Stau2 targets, with Sally Temple's group identifying genome-wide targets in the entire cortex and Freda Miller's group focussing on specific candidates. Gene ontology (GO) analysis showed Stau2 RNA targets were enriched in transcripts encoding regulators of cell-cycle exit and cilia.³³ Of note, several of these targets, *Prox1*, *Bbs2* and *Trim32*, are asymmetrically localized in a Stau2-dependent fashion. Altogether, these results suggest that Stau2 plays a preponderant role in the selective transmission of pro-differentiation mRNAs in progeny. In parallel, given other RNA regulatory roles of Stau2, the RBP may additionally repress translation of pro-differentiation mRNAs in NSCs. The authors speculated that this translational repression would be relieved in differentiated daughter cells.

These studies indicate that RNA localization may serve as a cell fate determinant to help two daughter cells adopt different fates. Utilizing the "RNA medium" to segregate cell fate determinants represents a certain advantage. As translation is largely paused during mitosis⁸⁷ the inheritance of select mRNA molecules in daughter progeny might allow for the rapid and massive synthesis of this determinant immediately after completion of mitosis. This mechanism involves a multistep process which includes: (1) the production of an mRNA and its associated RBPs prior to mitosis, (2) the translational repression of this mRNA to a cellular region which will be specifically inherited by one daughter cell, and (4) the derepression of translation following mitosis. Future studies and identification of

asymmetrically localized mRNAs and RBPs in mitotic neural progenitors will help define whether this mechanism is broadly used for cell fate determination in the mammalian cortex.

CONCLUSION

This review highlighted key players in posttranscriptional RNA regulation with fundamental roles in corticogenesis. One theme that emerges from this review is that we have just scratched the surface in terms of a comprehensive understanding of how RBPs influence cortical development and which RBPs are important. A second theme that emerges is that virtually all aspects of posttranscriptional regulation are implicated in corticogenesis. Many fundamental questions now remain to be answered. How is RNA regulation coordinated within rapidly dividing populations across stages of embryonic development? What additional RBPs influence corticogenesis, how do they do so, and what are their key targets? What role does RNA stability play in modulating cell fate choices in the developing brain? Answering these questions in a complex tissue such as the embryonic mammalian cortex is challenging and will require multidisciplinary approaches encompassing bioinformatics, biochemistry, and genetics.

Identifying RNA targets for RBPs is critical to gain a mechanistic understanding of how these RBPs help shape the developing brain. Techniques in RNA immunoprecipitation have been critical for uncovering RNA targets for many RBPs within immortalized cells. With variations on RIP approaches, such as HITs-CLIP, it has now become possible to identify RNA targets within whole mouse brains. However, as noted throughout this review, approaches using entire tissue only give a superficial understanding of RBP targets, given the heterogeneous nature of the developing brain over time and space. Future studies which couple optimized RNA immunoprecipitation approaches with single cell resolution will be ideal. These will inform our understanding of how RBPs function in progenitors versus postmitotic neurons, and in early versus late development. Moreover, the use of Ribotagsequencing approaches or ribosomal profiling within neural stem cells will be extremely valuable to assess genome-wide translation profiles for RBPs. On an individual transcript basis, the direct visualization of RNA targets in situ is now also possible, using singlemolecule FISH probes. Using the MS2-tagging approach, one can also now follow single RNA movements in the cell.⁷⁹ Similarly, live-imaging can be used to evaluate translational targets, by use of reporter constructs in which regulatory mRNA sequences are tethered to photo-convertible protein, such as Kaede or Dendra.⁸⁸

Over 800 RBPs have now been bioinformatically and empirically identified, and among these many have annotated expression in the developing cerebral cortex.^{34,35,89} Yet it remains an outstanding question as to which of these RBPs are critical. These questions can be addressed using traditional genetics, CRISPR/Cas9 approaches, or *in utero* electroporation, the latter of which allows one to rapidly manipulate gene expression within the developing brain. Moreover, the ability to utilize primary cell culture of progenitors and *ex vivo* embryonic brain slice culture provide tools that make this developing organ amenable to testing candidates involved in posttranscriptional regulation. The future is exciting for RNA regulation in corticogenesis as the advent of new technologies will lead to great advances in this field of research.

ACKNOWLEDGMENTS

We thank members of the Silver lab for helpful discussions. Work in this lab is supported by an R01NS0830897 (to DLS). We apologize to all those whose work we could not include in this review.

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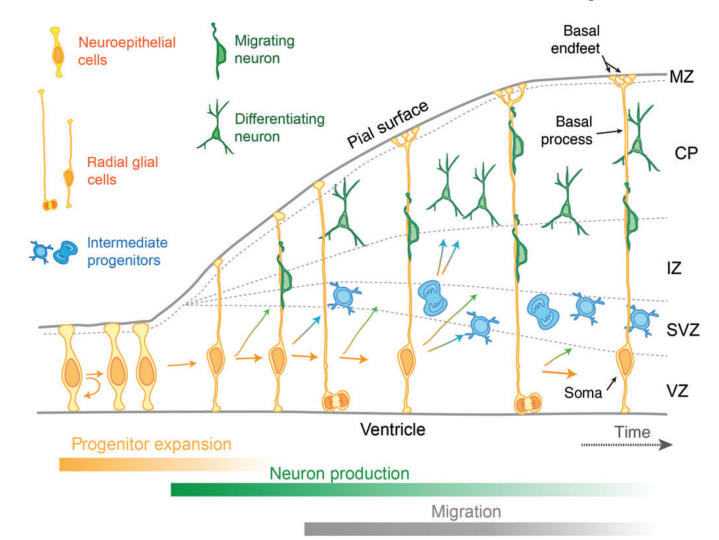


FIGURE 1.

Schematic representation of cortical development. Shown are three different progenitor populations (neuroepithelial cells, radial glial cells, and intermediate progenitors) and neurons (both migrating and differentiating). Progenitors residing within the VZ undergo self-renewal divisions to generate new progenitors (curved arrow) as well as divisions to generate either neurons or progenitors (straight arrows). As corticogenesis proceeds, progenitors initially expand their population, shift to neuron, and intermediate progenitor production. Intermediate progenitors within the SVZ also generate neurons. Neurons migrate through the IZ to the CP to form layers of the cerebral cortex. MZ, marginal zone; CP, cortical plate; IZ, intermediate zone; SVZ, sub-ventricular zone; VZ, ventricular zone

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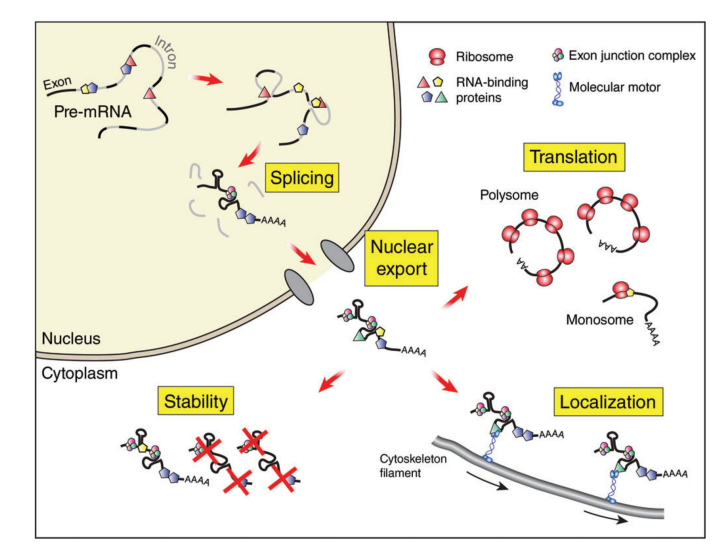


FIGURE 2.

Cartoon depicting various stages of mRNA life cycle when RBPs function. Different stages of posttranscriptional regulation are shown along with their nuclear-cytoplasmic location. This review discusses roles for RNA-binding proteins (shown as geometric shapes) in these various aspects of mRNA metabolism.

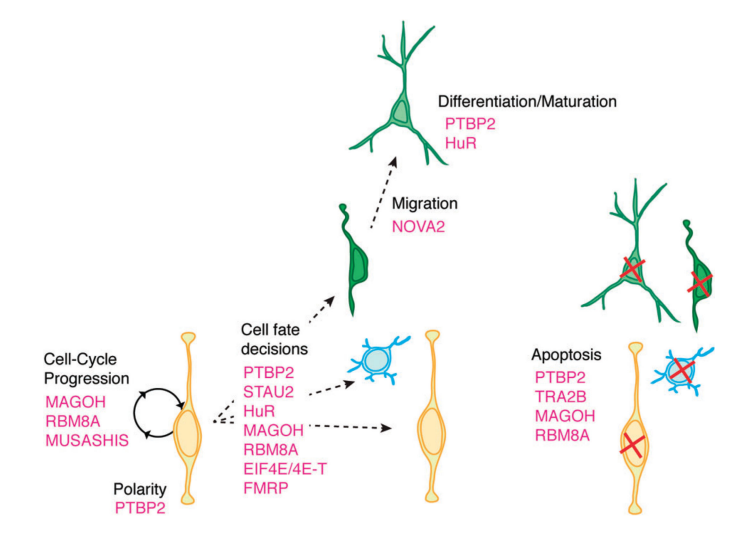


FIGURE 3.

Summary of known RNA-binding proteins and the aspects of corticogenesis they regulate. Different aspects of neural progenitor function (cell cycle progression, cell fate decision, apoptosis) and neuronal function (migration, differentiation, maturation, apoptosis) are indicated along with the RBPs discussed in this review.

TABLE 1

RNA-binding Proteins Required for Corticogenesis

RNA-binding Protein	Cortical Function	Cortical Expression	RNA Function	Key RNA Targets	Associated Neuro-diseases	References
NOVA2	Neuronal migration	Neuron specific	Alternative splicing	DabI	Paraneoplastic neurologic disorder	12,13
TRA2B	Neuronal and progenitor survival	Progenitors and neurons	SR pre-mRNA splicing	ć		14,15
PTBP2	Progenitor polarity, proliferation, neuronal maturation	Neuron specific	Alternative splicing, exon inclusion	Psd-95		16–18
MAGOH	Progenitor proliferation	Ubiquitous progenitor enriched	splicing, RNA localization NMD, translation	LisI		19
RBM8A	Progenitor proliferation	Ubiquitous progenitor enriched	splicing, RNA localization, NMD, translation	ć	TAR syndrome	20
HuR	Progenitor proliferation, neuronal identify and maturation	Neuron specific	Translation	DliJ		21,22
MSI1,2	Progenitor proliferation	Ubiquitous, enriched in progenitors	Translation	Numb, Jag1, Prfpf3, Kirrel3, Rbm22, Dhx37		23–25
FMRP	Progenitor maintenance	Progenitors and neurons	Translation	Pfn1, Cdh2, NOS1	Fragile X	26–30
Eif4E/4E-T	Progenitor maintenance	Progenitors and neurons	Translation	Neurog1, Neurog2, Neurod1		31
STAU2	Radial glia maintenance	Progenitors and neurons	RNA localization	Prox1, Bsb2, Trim32		32,33

TABLE 2

Genomic Studies Highlighting Alternative Splicing in Cortical Development

Stage	Organism	Analysis	References
Mid-gestation	Human	Spatial differences between brain structures	37
E14.5	Mouse	Spatial differences in cortical layers(VZ/SVZ vs IZ vs CP)	36
E15.5-P1	Mouse	Temporal and spatial differences in sorted excitatory neurons: callosal, subcerebral, corticothalamic/subplate neurons	40
E16-P30	Mouse	Temporal differences in in vivo cerebral cortices	41
In vitro	Human	Temporal differences in in vitro differentiated ES cells	42