Cytoplasmic RNA-Binding Proteins and the Control of Complex Brain Function

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The formation and maintenance of neural circuits in the mammal central nervous system (CNS) require the coordinated expression of genes not just at the transcriptional level, but at the translational level as well. Recent evidence shows that regulated messenger RNA (mRNA) translation is necessary for certain forms of synaptic plasticity, the cellular basis of learning and memory. In addition, regulated translation helps guide axonal growth cones to their targets on other neurons or at the neuromuscular junction. Several neurologic syndromes have been correlated with and indeed may be caused by aberrant translation; one important example is the fragile X mental retardation syndrome. Although translation in the CNS is regulated by multiple mechanisms and factors, we focus this review on regulatory mRNA-binding proteins with particular emphasis on fragile X mental retardation protein (FMRP) and cytoplasmic polyadenylation element binding (CPEB) because they have been shown to be at the nexus of translational control and brain function in health and disease.

The integrated circuit that is the mammalian CNS controls complex cognitive processes such as learning, memory, and behavior. Synapses, the points of communication between neurons, are essential for closing this circuit. They are morphologically dynamic structures that respond to neurotransmitter stimulation by increasing or decreasing the strength of their response, a phenomenon referred to as synaptic plasticity. In the most basic sense, synapses are composed of pre- and postsynaptic compartments; presynaptic boutons are axonal substructures that transmit signals, whereas postsynaptic domains primarily occur on the dendrites and cell bodies of the neurons that receive these signals. The adult presynaptic compartment generally lacks identifiable polysomes, suggesting that translation in this compartment is unlikely to play a substantial role in mature neuronal function. However, lack of detection of polysomes by electron microscopy is not proof of their absence, and indeed if only a few ribosomes are engaged with each messenger RNA (mRNA), then ultrastructural analysis may underestimate the translational capacity of this compartment. On the other hand, the postsynaptic compartment clearly contains polysomes, indicating that translation in this region could be important for higher-order brain activity (Steward and Levy 1982). Indeed, some forms of synaptic
plasticity, which comprise the underlying cellular basis of learning and memory, almost certainly require mRNA translation in the postsynaptic region (reviewed by Kandel 2001; Sutton and Schuman 2006; Richter and Klann 2009). Moreover, this translation is “local” because at least with Schaffer collateral CA1 neurons of the hippocampus, it takes place in the dendritic layer even when it is severed from the nucleus-containing cell body (Kang and Schuman 1996; Huber et al. 2000). Translational control and RNA localization in neurons have been reviewed by a number of investigators (Holt and Bullock 2009; Wang et al. 2010; Doyle and Kiebler 2011); here, we focus on RNA-binding proteins as regulators of translation and complex brain function.

TRANSLATIONAL REGULATION IN NEURONS

It is now accepted wisdom that memory consolidation requires protein synthesis during a defined period relative to the training stimulus (Flexner et al. 1962; Flexner and Flexner 1968; Sutton and Schuman 2006). The foundation of this consolidation requires long-term changes in synaptic efficacy, measured electrophysiologically as long-term potentiation (LTP) or long-term depression (LTD), or ultrastructurally as shape changes in dendritic spines (Ostroff et al. 2002, 2010; Mishchenko et al. 2010; Dent et al. 2011). Before discussing particular mechanisms of translational control by RNA-binding proteins in neurons, however, synaptic plasticity should be placed into a molecular and cellular context that addresses not only how translation is regulated, but why. Consider that a typical neuron in the CNS may have thousands of synaptic inputs, yet when stimulated, the neuron can distinguish between experienced (stimulated) and naive (unstimulated) synapses. How it does so has been a subject of intense investigation, but it is generally thought that the neuron “tags” the stimulated synapses, which results in changes in synaptic efficacy (Frey and Morris 1997). The nature of the tag may be complex, but at least in some circumstances, it involves de novo translation in dendrites. Thus, cellular memory, which when placed within a complex circuitry forms the basis for organismal memory, requires regulated translation. Moreover, certain stimulation protocols elicit changes in the synthesis of specific proteins (Scheetz et al. 2000), suggesting that general translational control mechanisms may not be sufficient for changes in plasticity. For reviews of the signaling pathways leading to changes in translation, see recent articles by Costa-Mattioli et al. (2009) and Hoeffer and Klann (2010).

FMRP AND THE FRAGILE X SYNDROME

Although the brain contains a multitude of mRNA-binding proteins, only a few have been shown to be required for proper neurologic function. Perhaps the most well known of these proteins is fragile X mental retardation protein (FMRP), the product of FMR1, the fragile X mental retardation syndrome gene. Individuals with the fragile X syndrome display a range of afflictions including epileptic-like cognitive deficits, autistic behaviors, epileptic-like seizures in childhood, and morphological anomalies such as elongated faces and large ears (Hagerman and Hagerman 2002; Penagarikano et al. 2007; Hagerman et al. 2009; Santoro et al. 2011). The syndrome is caused by a CGG triplet repeat expansion in the 5′ UTR of the FMR1 gene, resulting in abnormal DNA methylation and transcriptional silencing (Pieretti et al. 1991; Verkerk et al. 1991). FMRP is a complex RNA-binding protein that contains two KH (hnRNP K homology) domains and an RGG (arginine-glycine-glycine) box (Siomi et al. 1993), of which KH2 is perhaps the most critical for function (De Boulle et al. 1993; Zang et al. 2009). FMRP likely represses translation in dendrites as well as the cell body, yet when brain lysates are centrifuged through sucrose gradients, FMRP sediments with polysomes (Feng et al. 1997b; Khandjian et al. 2004; Stefani et al. 2004; Darnell et al. 2011), an unexpected association as translation is frequently repressed at the level of initiation and therefore most translational repressors are associated with mRNAs that are not polysome associated. The polyribosomal association of FMRP is dependent on the KH2
domain because a mutation in that region (I304N) abrogates RNA binding and polysome association (Feng et al. 1997a; Darnell et al. 2005a; Zang et al. 2009); as a consequence, expression of this nonfunctional protein elicits symptoms of the fragile X syndrome even when there is no CGG repeat (De Boulle et al. 1993). In addition, an I304N knockin mouse model also shows characteristics of the syndrome nearly identical to that of an FMRP knockout mouse (Zang et al. 2009). Taken together, these and other data indicate that the loss of FMRP function in regulating polysome-associated translational inhibition forms the molecular basis of the fragile X syndrome. It therefore follows that two keys for understanding the etiology of the syndrome lie with the identification of target mRNAs and the biochemical mechanism for their translational regulation.

The identification of mRNA targets and binding sites for RNA-binding proteins can be technically challenging processes. Two often-used approaches are in vitro RNA selection (Ellington and Szostak 1990) (also known as SELEX, selective enrichment of ligands by exponential enrichment [Irvine et al. 1991]) or coimmunoprecipitation of protein-RNA complexes followed by microarray analysis or RT-PCR identification of specific transcripts (RIP-Chip) (Tenenbaum et al. 2000). Although often successful, these methods have their limitations; SELEX-identified sequences do not take into account the in vivo milieu that can help determine sequence specificity and RIP-Chip has been criticized for its low stringency and a high rate of both false positives and false negatives (Mukherjee et al. 2011). To circumvent these issues, an in vivo UV cross-linking procedure (CLIP; when combined with high throughput sequencing, it is referred to as HITS-CLIP or CLIP-seq) has been devised that is highly specific with respect to both mRNA that is bound by a particular protein and the precise cis element that it recognizes (Ule et al. 2003; Licatalosi et al. 2008; Chi et al. 2009; Darnell et al. 2011; reviewed in Darnell 2010). HITS-CLIP uses UV irradiation to covalently link RNA-binding proteins and RNA, followed by stringent immunoprecipitation of the protein of interest, limited RNase digestion to reduce the size of the RNA that is “CLIPed,” addition of linkers, and high-throughput sequencing (Fig. 1). The details of the procedure have been described elsewhere (Ule et al. 2005; Jensen and Darnell 2008), as has an outline of the advantages this method has over other techniques (Darnell 2010; Zhang and Darnell 2011). Suffice it to say that CLIP and subsequent modifications (Granneman et al. 2009; Hafner et al. 2010; Konig et al. 2010) have now been widely used to analyze several RNA-binding proteins from a number of tissues (Darnell 2010).

FMRP offers a particularly illustrative example of the power of CLIP and the results that can be obtained. Previously, in vitro SELEX analysis of FMRP had shown that the RGG box bound a G-quadruplex structure (Darnell et al. 2001, 2004) and KH2 bound a “kissing complex” (Darnell et al. 2005b). Moreover, RIP-Chip analysis from mouse brain found that the protein interacted with $\approx 430$ mRNAs (Brown et al. 2001). Application of the HITS-CLIP technique to polysome-associated FMRP revealed that FMRP interacted with a large and specific set of mRNAs highly enriched for those encoding components of both the pre- and postsynaptic compartments (Darnell et al. 2011). Interestingly, about half of the RIP-Chip-identified FMRP target mRNAs could not be confirmed by in vivo UV cross-linking, possibly illustrating the issue of false positives in the RIP-Chip assay (as in any method, CLIP may also have false positives as well). Moreover, only about a quarter of the $\approx 840$ FMRP UV cross-linked mRNAs were found by the RIP-Chip approach, indicating a high false negative rate for RIP-Chip, likely because of their loss during immunoprecipitation owing to the lack of a covalent bond between protein and RNA. In addition and quite surprisingly, the preponderance of FMRP-binding sites (CLIP sequence tags) resided in mRNA coding sequences, and not in 5′ or 3′ UTRs as might be expected based on analogy with other RNA-binding proteins such as the neuronal Hu family (Darnell et al. 2011). Moreover, FMRP interacted with its target mRNAs with an even distribution throughout the coding sequence, similar to the distribution of ribosomes, with no discernible preference for sequence or structural
motifs (Fig. 2, the placement of the ribosomes on the mRNA is for illustrative purposes only and is not meant to convey a particular congre- gation at the 3′ end). These seemingly paradoxical observations lead one to ask how does FMRP associate with polysomes yet still repress translation, and how does FMRP bind certain mRNAs in an apparently cis element-independent manner? Finally, do the CLIP results give insight into the fragile X syndrome or autism spectrum disorders (ASD)?

The observation that FMRP represses translation (Laggerbauer et al. 2001; Li et al. 2001) but does so while associating with polysomes (Khandjian et al. 1996, 2004; Corbin et al. 1997; Feng et al. 1997a,b; Stefani et al. 2004) has...
given rise to the idea that it inhibits polypeptide elongation (ribosome transit) (Ceman et al. 2003) but much of the initial data supporting that idea was based on overexpression of FMRP, cultured cells, and the use of copy DNA (cDNA) reporter constructs. The identification and validation of a robust set of in vivo mRNA targets of FMRP, and an equally high-confidence set of nontarget mRNAs as controls, allowed evaluation of this hypothesis in a physiologically relevant setting (Darnell et al. 2011). Darnell et al. devised an in vitro assay of translation using the same starting material used to measure FMRP mRNA binding (a polyribosomal extract from mouse brain) so that mRNA binding could be correlated with function with statistical significance. They found that FMRP target mRNAs were associated with stalled ribosomes that were resistant to runoff, whereas mRNAs that were not bound to FMRP (as assessed by HITS-CLIP) showed a typical run-off profile and shifted from the polysome region to lighter fractions of the sucrose gradient. Although the mechanism by which FMRP stalls ribosomes remains to be determined, it seems clear that this is the primary mechanism by which FMRP inhibits translation (illustrated in Fig. 2).

Presumably related to ribosome stalling is the fact that FMRP interacts with coding sequences. Does FMRP act as a “roadblock” for transiting...
ribosomes as has been suggested for ash1p mRNA in yeast (Chartrand et al. 2002)? Does it act in a manner analogous to SRP, which causes reversible ribosome stalling on mRNAs encoding secreted or membrane-associated proteins (Wolin and Walter 1988)? Perhaps FMRP blocks peptide elongation through interactions with eukaryotic elongation factor 2 (eEF2) (Sutton et al. 2007; Park et al. 2008). Much remains to be determined about FMRP’s mechanism of action and whether its apparently reversible action to repress translation is mediated by its phosphorylation state (Ceman et al. 2003), turnover (Hou et al. 2006), or another mechanism.

Finally, the set of mRNA substrates to which FMRP cross-links begins to offer some insight into the etiology of the fragile X syndrome specifically, and ASDs more generally. For example, of the 842 FMRP target mRNAs, 28 have also been linked to ASDs including neuroligin 3, neurexin 1, shank3, PTEN, TSC2, and neurofibromatosis 1, suggesting that alterations in the functional levels of these proteins in neurons may underlie the common symptoms in fragile X syndrome and autism (Darnell et al. 2011). Other FMRP target mRNAs whose encoded proteins are implicated in ASDs include those involved in the ERK and mTOR pathways, NMDA receptor complexes, regulators of small GTPases, and cell–cell adhesion molecules (Kelleher and Bear 2008; Hoeffer and Klann 2010). The FMRP CLIP data have revealed other potential insights into FMRP function. Consider that about one-third of the FMRP CLIP targets encode presynaptic proteins (Darnell et al. 2011). Indeed, a presynaptic function for FMRP has been suggested by localization studies that have found FMRP in axons and growth cones (Feng et al. 1997b; Antar et al. 2006; Christie et al. 2009) as well as the characterization of axonal growth cone motility (Antar et al. 2006; Li et al. 2009), elongation (Tessier and Broadie 2008), and pathfinding (Michel et al. 2004) defects in both fly and mouse models of fragile X syndrome. In addition, there is experimental support for a presynaptic role in synapse formation and the establishment of circuitry (Zhang et al. 2001b; Hanson and Madison 2007; Bureau et al. 2008; Gibson et al. 2008). Recent studies on the role of the Aplysia FMRP homolog (ApFMRP) in sensory to motor neuron synaptic plasticity support both a pre- and postsynaptic role for FMRP in regulating protein synthesis in response to synaptic stimulation (Till et al. 2011). A fruitful area for further research is to connect these observations with the set of presynaptic mRNAs whose translation is regulated by FMRP (Darnell et al. 2011).

CPEB
The cytoplasmic polyadenylation element-binding protein CPEB is an mRNA-specific translational control factor that was initially identified for its role in regulating cytoplasmic polyadenylation during Xenopus oocyte maturation (reviewed in Mendez and Richter 2001). CPEB has two RNA-recognition domains (RRM motifs) and two zinc-finger motifs (Hake and Richter 1994; Hake et al. 1998), and by binding to the 3′ UTR cytoplasmic polyadenylation element (CPE) found in specific mRNAs, it recruits a number of interacting proteins to modulate poly(A) tail length and as a result, translation (Richter 2007). CPEB activity begins in the nucleus, where it binds the CPE of pre-mRNAs, which like most nuclear pre-mRNAs, probably have long poly(A) tails (Lin et al. 2010). Following export to the cytoplasm, CPEB associates with a number of factors including Gld-2, a noncanonical poly(A) polymerase, and PARN, a poly(A) ribonuclease. PARN is the more active of the enzymes and thus shortens the poly(A) tail to usually \( \approx 20–40 \) nucleotides. Hormonal stimulation leads to CPEB phosphorylation, which expels PARN from the RNP complex; as a consequence, the poly(A) tail is elongated by Gld2 (Barnard et al. 2004; Kim and Richter 2006, 2007). The poly(A) tail is then bound by poly(A)-binding protein (PABP), which in turn, helps recruit the initiation complex to the 5′ end of the mRNA. Indeed in a general sense, genome-wide analysis has shown that poly(A) tail length is positively correlated with both the ribosome density on a transcript and the degree of association of PABP, providing additional support for the model that poly(A) tail length regulates translation through PABP.
and the initiation complex, primarily eIF4G (Halbeisen et al. 2008).

CPEB also represses translation by recruiting specific eIF4E-binding proteins (4E-BPs, neuroguidin in neurons, maskin in oocytes) to the mRNA (Stebbins-Boaz et al. 1999; Richter and Sonenberg 2005; Jung et al. 2006). 4E-BPs repress translation by binding eIF4E, the cap-binding protein, which prevents eIF4E from associating with eIF4G and thus inhibits initiation. Neuroguidin localizes to puncta in dendrites and growth cones, and is frequently seen at the very tip of filopodia and at the leading edges of growth cones. Knockdown of the neuroguidin homolog in Xenopus causes a failure of neural tube closure and inhibition of neural crest cell migration, suggesting an important role for CPEB- and neuroguidin-mediated translational repression in neurons during development (Jung et al. 2006).

Vertebrates contain four CPEB paralogs; CPEB1 described above may be functionally distinct from CPEB proteins 2–4. Inhibition of CPEB1 activity, through expression of dominant–negative forms of the protein or in Cpeb1 knockout (KO) mice, leads to defects in learning, memory, synaptic plasticity, dendritic arborization, and neuronal circuit formation (Alarcon et al. 2004; Berger-Sweeney et al. 2006; McEvoy et al. 2007; Bestman and Cline 2008). CPEB1 inhibits translation of its target mRNAs until glutamatergic activation (Wells et al. 2001) stimulates its phosphorylation by either Aurora kinase A (Mendez et al. 2000a,b; Huang et al. 2002) or Camk2a (Atkins et al. 2004, 2005), resulting in increased mRNA polyadenylation and translation at synapses. The strongest evidence for participation of CPEB1 in local translation is its localization to postsynaptic sites in dendrites (Wu et al. 1998) and enrichment of phospho-CPEB1 in a postsynaptic density (PSD) fraction purified from rat neurons (Atkins et al. 2004), although it is not entirely clear that the antibodies used did not cross-react with either the CPEB2-4s or other phosphoproteins because a Cpeb1 KO was not available as a negative control at that time.

Recently it was suggested that CPEB1 may also regulate local translation in hippocampal growth cones in response to neurotrophin stimulation through regulation of CPE sites in β-catenin mRNA (Ctnnb1) (Kundel et al. 2009). This raises an interesting issue in the context of a related study by Holt and colleagues, also addressing whether CPEB regulates local translation in the growth cone (Lin et al. 2009). In the latter study both the CPE and cytoplasmic polyadenylation of mRNAs were required for translation-dependent chemotropic responses, but CPEB1 itself was not. Because other RNA-binding proteins can potentially bind the CPE and regulate cytoplasmic polyadenylation (Slevin et al. 2007), studies using overexpression of the CPEB RNA-binding domain (RBD) to inhibit CPEB activity may also block CPE binding by other RNABPs. The CPE, with a consensus of $U_4-AUU$, is similar to other U-rich or UA-rich binding sites for the large family of ARE-binding proteins, including Hu (discussed below), and, in fact, the Xenopus homolog of Hu, ElrA, has been reported to bind the CPE (Slevin et al. 2007). The extent to which mammalian neuronal Hu isoforms may interact with CPEB1 to control translation is an interesting question for further study.

In sum, although it is likely that CPEB1 fulfills an important role in regulating neuronal translation it is not fully established that CPEB1 is responsible for local translational control underlying long-term synaptic plasticity changes. CPEB2–4 are also expressed in neurons and there is significant data from Aplysia and fly homologs to suggest that they may also regulate local translation. CPEB2–4 differ from CPEB1 in that they do not bind the CPE and do not regulate polyadenylation but instead recognize a different RNA consensus sequence (Huang et al. 2006). They also lack the Aurora kinase A phosphorylation site of CPEB1 (Theis et al. 2003). They have been shown to regulate translation in an N-methyl-D-aspartate receptor (NMDAR)–dependent manner in neurons, through a mechanism independent of CPE binding and polyadenylation (Huang et al. 2006).

The Aplysia CPEB homolog, ApCPEB, is neuron specific and most similar to CPEB3 (Liu and Schwartz 2003; Si et al. 2003a; Theis et al. 2003). Like CPEB3 and 4 in mouse, ApCPEB is induced in response to synaptic
activation (Theis et al. 2003) and is required for long-term facilitation at the activated synapse, strengthening the model that “type 2” neuronal CPEBs regulate protein synthesis-dependent forms of long-term synaptic plasticity. Owing to an amino-terminal glutamine-rich domain, ApCPEB has the additional property of forming prionlike particles that may be related to synaptic changes underlying long-term memory storage or form a synaptic “tag” marking that synapse (Darnell 2003; Si et al. 2003b, 2010). Notably, loss of function of the fly homolog of CPEB1 (Orb) has phenotypes in the fly oocyte (Lantz et al. 1994) but has not been linked to a neuronal defect. However, the CPEB2-4 homolog, Orb2, expressed predominantly in neurons and also containing a prionlike domain, is necessary for some forms of long-term memory (LTM). Intriguingly, its effects on LTM are dependent on the prionlike domain (Keleman et al. 2007). Taken together, results from Aplysia and flies suggest that elucidation of the mRNA targets and functions of CPEB2-4 in addition to CPEB1 will clarify mechanisms underlying some forms of LTM.

PUMILIO
The regulation of cytoplasmic polyadenylation is an important point of translational control for another family of RNABPs, the pumilio (pum) or PUF family (Wickens et al. 2002; Quenault et al. 2011). PUF proteins harbor a conserved Pumilio homology domain that binds RNA, and the proteins regulate cytoplasmic poly(A) length and may compete or cooperate with CPEB1. Like CPEB1, Pumilio regulates gene expression through two mechanisms, in this case by (1) affecting the stability of target mRNAs through recruitment of a deadenylase complex to shorten poly(A) tails and promote mRNA turnover (reviewed in Quenault et al. 2011) and (2) repressing translation directly. A carboxy-terminal RBD in Pumilio, first described by Zamore and colleagues (1997), recognizes the 16 nt “nanos response element” (NRE) in the 3′ UTR of hunchback mRNA in fly embryos (Wharton and Struhl 1991). Pumilio recruits two other proteins: Nanos, which is also involved in RNA localization, and Brat, which binds the eIF4E-like cap-binding protein d4EHP (Cho et al. 2006). Unlike eIF4E, d4EHP does not interact with delf4G; consequently initiation does not occur because without eIF4G, the 4S ribosomal subunit is not recruited to the mRNA (Cho et al. 2006). Pumilio also has been shown to have cap-binding activity itself and so may compete with eIF4E to block initiation (Cao et al. 2010). Pumilio activates translation in some cases through cooperative binding with CPEB1 on transcripts containing both NRE and CPE elements, and may stabilize CPEB binding, but translational activation does not appear to involve altered polyadenylation of target mRNAs (Pique et al. 2008). Pumilio has been found to associate with a Nanos homolog and CPEB in Xenopus (Nakahata et al. 2001), supporting the conservation of this mechanism.

There are two human homologs, Pum1 and 2, present in the brain. To date, neither has been linked to a human cognitive disease, and there are no reports of neuronal defects in KO mice to date. Nonetheless, compelling evidence exists for an important role in mammalian neuronal function (reviewed in Baines 2005). Loss of Pumilio leads to defective LTM in flies (Dubnau et al. 2003), likely owing to its effects on dendrite morphogenesis as well as synapse growth and function (Mee et al. 2004; Menon et al. 2004; Ye et al. 2004). Pum2 is present in RNPs in the cell body and dendrites of rodent hippocampal neurons (Vessey et al. 2006). Studies using bidirectional manipulation of Pum2 levels show reciprocal changes in dendrite outgrowth, the density and morphology of dendritic spines, and the frequency of miniature excitatory post-synaptic currents (mEPSCs) (Vessey et al. 2010), revealing that Pum2 negatively regulates all three phenotypes. The observed increased mEPSC frequency appears to be owing to an increased number of excitatory synapses on dendritic shafts in the absence of Pum2 (Vessey et al. 2010).

Zip Code Binding Protein
Zip code binding protein 1 (ZBP1) is an interesting example of a neuronal RNA-binding protein recognized for its role in mRNA localization.
that also represses translation. ZBP1 harbors 4 KH-type RBDs. The homologs of chicken ZBP1 (mammalian IMP1-3 or IGF2BP1-3) are onco-fetal proteins that are highly expressed during development, absent postnatally, and are reexpressed in a high percentage of tumors (Tessier et al. 2004). Nonetheless, ZBP1 appears to have an important role in developing neurons. It is present in granules in the somatodendritic compartment together with β-actin mRNA, actb, through interactions with a 54-nucleotide binding site (the “zip code”) in the 3’ UTR (Zhang et al. 2001a). These granules contain components of the exon junction complex, indicating that they are translationally repressed on exiting the nucleus (Jonson et al. 2007). In primary neurons, the levels of ZBP1 in growth cones (Zhang et al. 2001a), dendrites, and spines (Tiruchinapalli et al. 2003) is positively correlated with neuronal activity. Significantly, knockdown of ZBP1 in cultured neurons eliminates a BDNF-stimulated increase in actin-rich dendritic spines (Eom et al. 2003). Knockdown or overexpression of ZBP1 in cultured neurons was found to decrease complexity of the dendritic arbor and this was dependent on the ZBP1 RBDs (Perycz et al. 2011). No effect was found on established arbors, suggesting an important role for ZBP1 during the critical period of dendritogenesis (Perycz et al. 2011).

Direct translational control by ZBP1 in neurons was shown by Huttelmaier and colleagues who found that ZBP1 repressed initiation on neuronal actb mRNA by binding to the 3’ UTR zip-code sequence and interfering with 80S complex formation (Huttelmaier et al. 2005). Mechanistic studies revealed that phosphorylation of ZBP1 by Src kinase on Y396 decreased binding to the zip code and allowed initiation, resulting in local synthesis of β-actin (Huttelmaier et al. 2005). This mechanism was also reported to regulate local translation of actb in growth cones to allow turning in response to brain-derived neurotrophic factor (BDNF) (Sasaki et al. 2010). Furthermore, nonphosphorylatable Y396F-ZBP1 was unable to rescue the ZBP1 knockdown effect on dendritic arborization (Perycz et al. 2011). ZBP1 has also been reported to increase internal ribosome entry site (IRES) -dependent initiation of the hepatitis C virus RNA by interacting with both 5’ and 3’ UTRs (Weinlich et al. 2009); however, a function to regulate IRES-dependent initiation in neurons has not been established.

**CAPRIN**

Caprins1 and 2 (also known as RNG105 and RNG140) are among the newest additions to the list of potential translational regulators in neurons. Caprins bind RNA through an amino-terminal coiled-coil motif and a carboxy-terminal RGG box. Caprin1 (Grill et al. 2004) associates with polyribosomes in rat cortex (Angenstein et al. 2005) and is a component of the same ribosome-containing granules as the fragile X mental retardation autosomal homolog proteins FXRPs (Shiina et al. 2005; Shiina and Tokunaga 2010), suggesting a role in translational regulation similar to that of the FMRP family. Knockdown of caprin1 or caprin2 in cultured mouse neurons causes a reduction in the number, length, and branching of dendrites (Shiina and Tokunaga 2010) and studies on primary neuronal cultures from caprin1 KO mice show defects in synapse development (Shiina et al. 2010). BDNF treatment caused the dissociation of caprin1 and its mRNA targets from these granules into polyribosomes, suggesting that neural activity relieves translational repression imposed by caprin1 (Shiina et al. 2005). Caprin1 and caprin2 localize to different granules in dendrites and so appear to have distinct functions in neurons, although both appear to be involved in dendrite and spine development.

The caprins nonspecifically repress translation in vitro translation systems and globally repress translation when overexpressed in vivo. These findings might suggest that the caprins repress translation nonspecifically. However, loss of function of caprin does not increase global translation rates, raising a more general issue regarding overexpression of RNABPs in functional assays, which has been addressed in the context of caprin-induced stress granule (SG) formation (Solomon et al. 2007). SGs are the result of clustering of mRNAs, stalled pre-initiation complexes, and RNABPs caused by
phosphorylation of eIF2α or inhibition of the eIF4F complex (see Decker and Parker 2012). Caprins belong to a group of mRNA-binding proteins (including FMRP) that nucleate SGs when expressed at greater than physiological levels. Solomon et al. (2007) found that caprin overexpression caused eIF2α phosphorylation resulting in both global translational repression and the formation of SGs. Although it is possible that this might be related to the physiological role of caprins, it raises a red flag concerning the assignment of the role of translational repressor based solely on overexpression studies.

HU

The mammalian neuronal homologs of the Drosophila ELAV protein (Hu B, C, and D) have been reported to regulate mRNA stability (Bolognani and Perrone-Bizzozero 2008), translation (Fukao et al. 2009), and pre-mRNA splicing (Zhu et al. 2006) to regulate nervous system development and function (Okano and Darnell 1997; Akamatsu et al. 1999; reviewed by Antic and Keene 1997; Hinman and Lou 2008). ELAV is known to be essential for nervous system development and function in flies (Campos et al. 1985; Robinow et al. 1988). The mammalian homologs were first cloned as antigens of paraneoplastic antibodies in the human neurologic Hu syndrome (Szabo et al. 1991; Posner and Dalmu 1997) and studies using mice in which these proteins have been knocked out or overexpressed show a number of neuronal, learning, and memory phenotypes (Akamatsu et al. 2005; Bolognani et al. 2006, 2007a,b; Tanner et al. 2008). Hu has been consistently shown to be up-regulated in hippocampal neurons after contextual (Bolognani et al. 2004) or spatial learning tasks (Quattrone et al. 2001; Pascale et al. 2004) and after glutamate receptor activation (Tiruchinapalli et al. 2008). The roles of the neuronal Hu proteins in development, plasticity, and memory have been recently reviewed (Deschenes-Furry et al. 2006; Pascale et al. 2008).

Hu proteins have three RRM domains and bind to AU- and GU-rich elements in the 3′ UTRs of specific mRNA transcripts. These elements, also known as AREs, are also binding sites for a number of RNABPs that promote turnover of these mRNAs, including KSRP, TTP, AUF1, and BRF1. Hu is thought to bind the same or similar elements to protect ARE-containing transcripts from degradation. As a group, the ARE-binding proteins regulate mRNA stability in response to cell stimulation including stress, proliferative stimulation, immune signaling, and developmental signals.

Hu can directly regulate the translation of target mRNAs as well. Neuronal Hu’s are localized to somatic and dendritic granules containing ribosomes (Bolognani et al. 2004) and are polyribosome-associated in human medulloblastoma cells (Gao and Keene 1996), PC12 cells (Fukao et al. 2009), primary neurons (Tiruchinapalli et al. 2008), and brain (Bolognani et al. 2004; Darnell et al. 2009). Polyribosome-associated Hu levels increase after KCl stimulation (Tiruchinapalli et al. 2008) or learning (Bolognani et al. 2004), suggesting that Hu increases translation of its mRNA targets in response to neuronal activity by promoting initiation (Chen and Shyu 2009; Fukao et al. 2009) supported by the increased association of its mRNA targets with polyribosomes (Antic et al. 1999; Mazan-Mamczarz et al. 2003; Kawai et al. 2006; Galban et al. 2008). Stimulation of translation by HuD has been shown to be dependent on both the poly-A tail and 5′ cap and occurs through direct interaction of HuD with the helicase eIF4A (Fukao et al. 2009). This mechanism of facilitating initiation is critical for Hu-dependent neurite outgrowth in PC12 cells (Fukao et al. 2009). Hu has also been proposed to inhibit translation in some cases (reviewed in Hinman and Lou 2008).

CONCERTED FUNCTION OF TRANSLATIONAL REGULATORY PROTEINS IN NEURONS

In sum, intriguing evidence supports the idea that specific mRNA-binding proteins regulate translation that in turn mediates neuronal function including synaptic plasticity. Moreover, mounting evidence suggests that such regulation is likely to require the concerted action of
the translational regulators discussed in this review. Although identification of important neuronal translational regulatory proteins through genetic and biochemical approaches has been quite successful, identification of the mRNAs whose expression is controlled by these binding proteins remains a major hurdle limiting our understanding of the mechanisms involved. Application of HITS-CLIP and related techniques to FMRP, CPEB, Pumilio, ZBP, caprin, and Hu in brain, and their use in quantifying changes in RNA binding in response to activity, or in subcellular fractions such as polyribosomes or purifiable granules, or during development will likely lead to dramatic advances in our understanding of how these proteins fine tune the synthesis of key neuronal proteins.

**RELEVANCE TO THERAPY FOR HUMAN COGNITIVE AND BEHAVIORAL DISEASES**

One approach to modulating translation in neurons to ameliorate symptoms of diseases such as fragile X syndrome or some cases of autism, both thought to be caused by “runaway translation,” is to target the general signaling pathways that transduce engagement of cell-surface receptors to increased translation, including PI3K/Akt/mTOR and MEK/ERK (Kelleher and Bear 2008; Hoffer and Klann 2010). These pathways are already the subject of anticancer therapy research because the cell cycle uses similar pathways to transduce growth factor signals into translational output as do neurons (Guer廷 and Sabatini 2005; Lee and Blenis 2005; Sabatini 2006; Frost et al. 2009; Dowling et al. 2010; Livingstone et al. 2010; Silvera et al. 2010). Hopefully, very specific therapies may result from understanding binding sites and mechanisms of translational control at the molecular level, both of which can be addressed through HITS-CLIP “mapping” of physiologic RNA-binding protein:RNA interactions paired with appropriately designed mechanistic studies using endogenous mRNA targets. The majority of the mRNA translational regulatory proteins are thought to repress translation whether at the level of elongation, like FMRP, or initiation, like CPEB, pumilio, and ZBP. Replacement of the repression lost in disease might be achieved by delivery of a mimic of FMRP activity to stall ribosomes, for example. Furthermore, it’s possible that generally slowing translation with the off-target use of antibiotics might take the edge off of excessive translation and provide therapeutic benefit, as has been suggested for fragile X syndrome (Darnell et al. 2011).

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