Regulation of Pluripotency by RNA Binding Proteins

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Establishment, maintenance, and exit from pluripotency require precise coordination of a cell’s molecular machinery. Substantial headway has been made in deciphering many aspects of this elaborate system, particularly with respect to epigenetics, transcription, and noncoding RNAs. Less attention has been paid to posttranscriptional regulatory processes such as alternative splicing, RNA processing and modification, nuclear export, regulation of transcript stability, and translation. Here, we introduce the RNA binding proteins that enable the posttranscriptional regulation of gene expression, summarizing current and ongoing research on their roles at different regulatory points and discussing how they help script the fate of pluripotent stem cells.

Introduction
Embryonic stem cells (ESCs), which are derived from the inner cell mass of the mammalian blastocyst, are remarkable because they can propagate in vitro indefinitely while retaining both the molecular identity and the pluripotent properties of the peri-implantation epiblast. Consequently, ESCs provide a biologically relevant and experimentally tractable model system for studying regulators of cell fate and cell fate transitions in early development. Understanding the molecular mechanisms of ESC maintenance and differentiation is critically important not just scientifically but also clinically, because an improved knowledge of pluripotency and embryonic development will allow ESCs to be more effectively utilized as an in vitro platform for disease modeling, drug discovery, and tissue regeneration.

While the transcriptional, signaling, and epigenetic regulation of these cells have been the primary focus of research efforts in recent years (reviewed in Ng and Surani, 2011; Young, 2011; Watanabe et al., 2013), posttranscriptional and translational mechanisms of control remain relatively unexplored, despite evidence that they play a dominant role in driving ESC fate decisions. Indeed, posttranscriptional regulation has been reported to account for nearly 75% of the changes in protein levels after differentiation induced by knockdown of the transcription factor Nanog (Lu et al., 2009), and it was recently demonstrated that control over translational initiation by the eIF4e binding proteins dramatically influences the efficiency of reprogramming somatic cells to induced pluripotent stem cells (iPSCs) (Tahmasebi et al., 2014). The cell controls protein levels posttranscriptionally using a large collection of tools that includes noncoding RNAs and RNA binding proteins (RBPs). Recent work elucidating the functions of microRNAs (miRNAs) and long noncoding RNAs (lncRNAs) in ESCs has been comprehensively reviewed elsewhere (Greve et al., 2013; Ghosal et al., 2013; Wright and Ciosk, 2013; Flynn and Chang, 2014). Similarly, posttranslational regulation of protein levels through the addition of covalent modifications also has been discussed recently (Wang et al., 2014b). The purpose of this Review is specifically to address the roles of RBPs in the maintenance and differentiation of ESCs.

RBPs are responsible for every event in the life of an RNA molecule, including its capping, splicing, cleavage, nontemplated nucleotide addition, nucleotide editing, nuclear export, cellular localization, stability, and translation (Keene, 2007). Overall, little is known about RBPs: most are classified based on computationally predicted similarities to proteins with known RNA binding domains, and until recently, few of these predictions have been verified in a cellular context in vivo. The recent introduction of a technique termed “mRNA interactome capture,” which enables the identification of proteins bound to polyadenylated RNAs in vivo, has been a significant development for the field (Baltz et al., 2012; Castello et al., 2012). Using this method, several groups were able to create a comprehensive catalog of RBPs in different mammalian cells, including 555 RBPs in mouse ESCs (Kwon et al., 2013). However, the functions of these RBPs in ESCs and their changes in ESC differentiation have yet to be addressed. Indeed, the mechanism of action of only a small number of RBPs has been examined in any great detail in the context of pluripotency. Here, we summarize current knowledge of RBP contribution to posttranscriptional and translational regulation in ESCs, following the approximate order that each regulatory event would be encountered as a transcript born in the nucleus migrates into the cytoplasm and is translated into a polypeptide (diagrammed in Figure 1). Throughout, we also discuss potential directions of future inquiry that will allow us to more fully appreciate the scope of RBP-mediated posttranscriptional and translational regulation in pluripotency.

Alternative Splicing
Alternative splicing of mRNA transcripts is probably the best-studied area of RBP-driven posttranscriptional control in ESCs, both in terms of the maintenance of the pluripotent state as well as in the artificial induction of pluripotency in somatic cells. A number of key pluripotency factors have been shown to exist as multiple isoforms that vary in stability, function, and intracellular localization due to differences in either coding exons or untranslated regulatory sequences. For example, Oct4 has two isoforms: Oct4A is the canonical pluripotency transcription factor expressed in ESCs and embryonal carcinoma cells, while
Oct4B is a cytoplasmic protein with unknown functions expressed in nonpluripotent cells (Atlasi et al., 2009). Intriguingly, Oct4 was identified in the HeLa mRNA interactome (Castello et al., 2012), raising the possibility that certain isoforms of Oct4 could be acting as RBPs in particular cell types. Other important alternatively spliced genes affiliated with the ESC state include Sall4, which has two isoforms that play collaborative roles in maintaining pluripotency; Tcf3, whose two isoforms inhibit Nanog and Oct4 transcription to different degrees (reviewed in Cheong and Lufkin, 2011); Nanog, whose three isoforms maintain ESC pluripotency with varying efficacies (Das et al., 2011); and DNMT3B, whose ~40 isoforms are each differentially expressed in pluripotent cells and various distinct tissue types (Go-palakrishnan et al., 2009). An especially well-developed story is that of Foxp1, which possesses several isoforms, including one that is uniquely expressed in ESCs. This particular transcript contains an exon variant that encodes an altered forkhead domain, yielding an ESC-specific Foxp1 with a DNA binding motif different from the canonical Foxp1 consensus motif. Demonstrating the dramatic impact of this change in sequence specificity, only the ESC form of Foxp1 can stimulate the transcription of pluripotency genes and repress ESC differentiation genes (Gabut et al., 2011). Thus, this interesting example shows how alternative splicing can be an integral part of the switch that regulates the choice between self-renewal and differentiation.

Transcript splicing is mediated by a core set of proteins and small nuclear RNAs that comprise the spliceosome, but additional RBPs are critical for dictating the spatial and temporal specificity of splicing events needed to drive tissue-specific expression of mRNA isoforms. Despite the obvious importance of this posttranscriptional phenomenon, the RBPs that enable the establishment and maintenance of the ESC splicing signature have only begun to be deciphered. Thus far, both positive (FOX2, SON, and SFRS2) and negative (MBNL1 and MBNL2) regulators of the ESC-specific splicing signature have been identified; the list of RBPs involved is sure to increase as research in this area expands.

In human ESCs (hESCs), FOX2 (RBM9 and FXH) promotes exon exclusion when bound to the upstream flanking intron and exon inclusion when bound to the downstream flanking intron, as demonstrated by crosslinking immunoprecipitation with high-throughput sequencing (CLIP-seq). Interestingly, many FOX2 targets are splicing factors as well, which suggests that FOX2 is an upstream master regulator of splicing regulators. FOX2 activity is critical for pluripotency, because depletion of the gene leads to hESC differentiation and death, thus highlighting the functional importance of active maintenance of the ESC splicing program (Yeo et al., 2009).

Another RBP critical for maintaining the hESC-specific complement of transcript variants is the spliceosome-associated factor SON. A genome-wide RNAi screen found that knockdown of SON caused hESC differentiation. Sequencing of wild-type and SON-depleted hESCs demonstrated that SON specifically targets short introns with GC-rich weak splice sites, and in the absence of SON, the transcripts of a number of key pluripotency genes like OCT4 and PRDM14 have aberrant inclusion of introns, leading to their degradation by nonsense-mediated decay (Lu et al., 2013). Like FOX2, then, SON helps create a splicing signature important for specifying and maintaining the pluripotent state.

A third splicing factor that positively regulates the self-renewing state of hESCs was described in a recent report (Lu et al., 2014). Sifting through transcriptome and proteome data comparing hESCs and hiPSCs with fibroblasts, these authors identified SFRS2 as the most enriched splicing factor in pluripotent cells and MBD2 as the most differentially spliced transcript.
This methyl-CpG binding protein exists as three isoforms (MBD2a–MBD2c), with MBD2c enriched in pluripotent stem cells and MBD2a in fibroblasts. Stable knockdown of SFRS2 in hESCs reduces expression of pluripotency genes and is associated with an isoform switch from MBD2c to MBD2a due to SFRS2 binding upstream of an exon unique to MBD2c. While overexpression of MBD2a promotes hESC differentiation, overexpression of MBD2c enhanced reprogramming of fibroblasts to hiPSCs. Interestingly, MBD2 had previously been identified as one of the miR-302 mRNA targets partially underlying the ability of this family of miRNAs to promote somatic cell reprogramming (Lee et al., 2013; Subramanyam et al., 2011). Here, the authors used this family of miRNAs to promote somatic cell reprogramming of MEFs. Splicing code analysis in human and mouse ESCs suggests that the presence of Mbnl and Mbd2 is one of the factors that determine the inclusion of repressive miRNA binding sites upstream of an exon unique to MBD2c. While overexpression of MBD2a promotes hESC differentiation, overexpression of MBD2c, suggesting that the effect of MBD2 on pluripotency is isoform dependent. In sum, SFRS2 provides an exciting example of how multiple regulatory pathways can cooperate to promote the pluripotent state.

In contrast to the positive regulators of ESC self-renewal described above, MBNL1 and MBNL2 repress ESC splicing patterns. Knockdown of either gene causes both human and mouse somatic cell lines to shift toward an ESC-like splicing pattern, including the acquisition of the ESC isoform of Foxp1. Moreover, depletion of Mbnl1 and Mbnl2 improved the efficiency of mouse somatic cell reprogramming of MEFs. Splicing code analysis in human and mouse ESCs suggests that the presence of Mbnl binding sites upstream of an exon is associated with its inclusion, while downstream binding is associated with exclusion—a pattern of binding opposite to that of FOX2 (Han et al., 2013).

Together, the examples of FOX2, SON, SFRS2, and MBNL1/2 reveal a very active role for RBPs that regulate splicing in both self-renewal and differentiation conditions. Determining how RBPs regulate splicing programs in cell fate choices encountered during the development of the three embryonic germ layers as well as the germline will be an important area of future pursuit.

**Alternative 3’UTR Cleavage and Polyadenylation**

In addition to varying the combination of coding exons that are included in an mRNA molecule, posttranscriptional regulatory machineries influence the processing of the 3’ ends of transcripts. Many mammalian genes contain more than one polyadenylation site that can be recognized for cleavage and addition of the poly(A) tail, and current estimates suggest that upward of 50%–70% of mammalian RNAs are subject to alternative polyadenylation (APA), leading to transcripts with varying 3’UTR lengths (Shepard et al., 2011; Derti et al., 2012). APA appears to be closely linked with cell state: for example, ESC differentiation and embryonic development is associated with 3’UTR lengthening (Ji et al., 2009; Shepard et al., 2011), while proliferation and somatic cell reprogramming are accompanied by 3’UTR shortening (Sandberg et al., 2008; Ji and Tian, 2009). This is thought to occur because the length of the 3’UTR can affect the availability of RBP and miRNA binding sites that are critical for determining transcript stability, localization, and translation. Indeed, there have been a few notable examples of how mRNA isoforms with shorter 3’UTRs can be more stable (Mayr and Bartel, 2009; Boutet et al., 2012) and more efficiently translated (Sandberg et al., 2008) than their longer counterparts due to differential inclusion of repressive miRNA binding sites. However, the biological roles of APA remain unclear at the global level, because sequencing studies, including one recently conducted in NIH 3T3 fibroblasts, have shown that long and short 3’UTR mRNA isoforms of the same genes have comparable stability and translation levels (Spies et al., 2013). These observations suggest, among other possibilities, that the effects of APA on transcript stability and translation may be specific to certain subsets of biologically important genes, while for the majority of transcripts, APA primarily serves some other regulatory purpose, such as intracellular localization (Spies et al., 2013).

Transcript cleavage and polyadenylation occur through the cooperation of several protein complexes: cleavage and polyadenylation specificity factor (CPSF) recognizes the polyadenylation signal (PAS) upstream of the cleavage site, while cleavage stimulating factor (CSTF) recognize downstream sequence elements (DSEs), and this leads to the recruitment of cleavage factors I and II, poly(A) binding proteins, and poly(A) polymerase (reviewed in Elkon et al., 2013). Interestingly, one of the protein subunits of CPSF, Fip1, was recently described to play a role in creating the APA profile of ESCs (Lackford et al., 2014). Consistent with its being important in pluripotency, Fip1 knockdown in ESCs promoted differentiation. Deep sequencing revealed that Fip1 depletion alters the APA profile of 374 genes, with most displaying a lengthening of their 3’UTR by shifting to the more distal PAS (proximal-to-distal shift [PtoD]). The genes showing PtoD were generally more highly expressed in self-renewing conditions than in differentiation conditions, and knockdown of a representative set partially recapitulated the Fip1 knockdown phenotype. Individual nucleotide crosslinking and immunoprecipitation followed by high-throughput sequencing (iCLIP-seq) showed that Fip1 binds U-rich sequences upstream of PAS sequences. Interestingly, when the PASs were located far apart, Fip1 bound upstream of the proximal PAS and promoted cleavage and polyadenylation at that site. However, when the PASs were close together, Fip1 promoted usage of the distal site. Given these data, the authors propose a model in which Fip1 promotes cleavage and polyadenylation at all PASs with upstream U-rich regions. However, when Fip1 is highly expressed, as it is in ESCs, it will bind and promote cleavage of the weaker proximal PAS, if there is a sufficient delay in transcription between the two PAS sites. This interesting model demonstrates how a general CPSF factor can regulate pluripotency by linking 3’UTR length with production of critical self-renewal factors. It will be important to investigate whether these results are unique to Fip1 or whether there are additional factors in ESCs that are critical for determining PAS usage in other sets of genes.

**RNA Postmodification**

Posttranscriptional RNA modification offers another exciting means for improved control over transcript levels and even increased proteomic diversity without the need to make changes to the genome. In regulatory regions, sequence alterations can affect transcript stability and translation efficiency, while in transcript coding regions, nucleotide changes can increase protein product diversity. RNA modifications can be separated into two types: the nontemplated addition of nucleotides and the chemical modification of templated bases. A well-established example of the former is the addition of nucleotides to the 3’ ends of RNA molecules. For instance, addition of uridines (uridylation) to pre-let-7 miRNA leads to its degradation in ESCs. In
particular, the 3′ terminal uridyl transferases (TUTases), Zcchc11 and Zcchc6, are directed by the RBP Lin28a to add a string of approximately 11 uridines to the pre-miRNA (Heo et al., 2008; Hagan et al., 2009). Oligouridylated pre-let-7 is then recognized by the DISL2 exoribonuclease, which catalyzes transcript degradation (Ustianenko et al., 2013). Interestingly, the addition of a single uridine has the opposite effect, promoting the processing of let-7 to its mature form (Heo et al., 2012). The remarkable number of pathways regulating let-7 activity befits its important role in promoting ESC differentiation and blocking reprogramming of somatic cells to iPSCs (Melton et al., 2010).

Although the existence of base modifications has been known for some time, their potential regulatory roles in mammalian cell fate decisions are just beginning to be uncovered. For example, the RNA editing enzyme Adar (Adar1) catalyzes the conversion of adenosine to inosine, which is subsequently read as guanosine by the translation machinery. Adar can target both mRNAs and dsRNA precursors of endo-siRNAs and miRNAs. In hESCs, ADAR was shown to drive A-to-I editing across the transcriptome, particularly in Alu repetitive elements. Furthermore, transient ADAR knockdown with an siRNA decreased A-to-I editing and led to an increase in the mRNA expression of genes associated with differentiation and development, although the ADAR-depleted ESCs did not exhibit significant morphological changes compared to wild-type (Osenberg et al., 2010). These results were based on the effects of a single transiently transfected siRNA, so it will be important to further examine the role of Adar by multiple different siRNAs, stable shRNA knockdown, and/or genetic deletion to rule out the possibility of off-target effects and to separate the effects of acute versus chronic loss of Adar. Moreover, the direct downstream consequences of A-to-I editing on targeted transcripts in ESCs remain to be elucidated. It is important to note that Adar has been found to have a number of different roles in other cellular systems. In a complex with Elavl1 (HuR), for example, Adar modulates mRNA stability (Wang et al., 2013a). Adar can also affect miRNA processing both negatively and positively, editing pri-miRNA transcripts to prevent Microprocessor cleavage (Wu et al., 2011) or dimerizing with Dicer in an RNA-dependent way to promote pre-miRNA cleavage (Ota et al., 2013). But because these studies were conducted in non-ESC cell lines, it would be meaningful to investigate whether these other diverse functions of Adar are also relevant in ESCs.

While ADAR-driven A-to-I base conversion is clearly important, methylation of adenosine to form N6-methyladenosine (m6A) is the most prevalent chemical modification of eukaryotic mRNA. Enriched at stop codons and 3′UTRs, the presence of m6A marks appears to influence many aspects of RNA metabolism. Indeed, recent evidence suggests that methylation at internal positions within transcripts is associated with transcript destabilization, while methylation at the transcriptional start site enhances translation (Schwartz et al., 2014). Biochemical studies recently revealed that these m6A marks are primarily deposited by the methyltransferases Mettl14 and Mettl3, which function together as a heterodimer (Liu et al., 2014). In mESCs, stable shRNA knockdown of either Mettl3 or Mettl14 greatly reduces m6A levels and impairs ESC self-renewal (Wang et al., 2014a). Interestingly, m6A-immunoprecipitation and sequencing (meRIP-seq) revealed that mRNAs of epigenetically poised developmental regulators (marked by H3K4me3 and H3K27me3 bivalent chromatin domains) are normally highly methylated, and these genes are upregulated in Mettl3- or Mettl14-depleted cells due to increased transcript half-life. These data suggest that m6A destabilizes transcripts. Further examining the mechanistic connection between methylation and RNA stability, the authors found that Mettl3/Mettl14 knockdown is associated with increased binding of the RBP HuR near m6A sites and leads to increased transcript stability, suggesting that HuR competes with Mettl3/Mettl14 and that their differential expression dynamically regulates RNA half-life. While it will be important to confirm these results in a genetic knockout system and further investigate the mechanism of methylated RNA destabilization, this study is intriguing for raising the idea that, like DNA methylation and chromatin modifications, “epigenetic” modification of RNA can be critical for determining cell identity.

Finally, the connections to HuR seen with both Adar- and Mettl3/Mettl14-mediated RNA editing are tantalizing, because HuR is also known to regulate transcript stability by either cooperating with or antagonizing miRNAs through a variety of different, context-dependent mechanisms. Additionally, HuR expression is itself modulated by miRNAs (reviewed in Simone and Keene, 2013). Together these studies exemplify how RNA modifications, RBPs, and miRNAs can function together to control steady-state mRNA expression levels.

Nuclear Export

In addition to directly altering the transcript sequences through alternative splicing, alternative polyadenylation, and nucleotide modifications, RNA fate can be influenced by the regulation of transport between the nucleus and cytoplasm, which controls protein levels by modulating access to the translation machinery. An exciting recent study has shown that such regulation can play a significant role in ESCs (Wang et al., 2013b). In an RNAi screen for factors that are required for ESC self-renewal, Hu and colleagues identified Thoc2 and Thoc5, two members of the THO complex, which provide an interface between nuclear transcription and RNA export. The authors show that knockdown of Thoc2 and Thoc5 does not alter overall transcript levels but rather leads to a nuclear accumulation of a subset of pluripotency mRNAs, including Nanog, Sox2, Klf4, and Esrrb. Immuno-precipitation followed by deep sequencing further showed that Thoc2 binds a subset of mRNAs in a Thoc5-dependent manner. Interestingly, Thoc5 is believed to be an adaptor protein, and its expression is downregulated with ESC differentiation. Overexpression of Thoc5 delays ESC differentiation, while knockdown promotes ESC differentiation and blocks somatic cell reprogramming. Therefore, the downregulation of Thoc5 that occurs during normal development appears to be critical for silencing parts of the pluripotency program and thus promoting differentiation. These findings uncover a new aspect of posttranscriptional control in pluripotency and may be just the first of many more examples to come of transcript-specific nuclear export influencing early embryonic cell fate decisions.

Regulation of mRNA Stability

Whether a molecule of RNA is stable or is quickly degraded in the cytoplasm governs, to a large extent, its ultimate protein...
expression levels. Transcript stability, which varies among cell types (Neff et al., 2012), is strongly influenced by RBP and miRNA binding, particularly at the 3’UTR. Three RBPs that have been implicated in this mode of regulation are Trim71 (miLin41), Pum1, and Brf1 (zfp36l1).

The role of Trim71 is noteworthy because it demonstrates an interconnection among multiple protein complexes. miRNAs, which typically regulate transcript stability and translation by binding to 3’UTRs, are critical for enabling the rapid proliferation of ESCs by shortening the G1 phase of the cell cycle (Wang et al., 2008, 2013c), and Gregory and colleagues found that Trim71 is important for facilitating this activity (Chang et al., 2012). Using immunoprecipitation followed by mass spectrometry, they found that Trim71 binds to miRNA-containing Ago2 complexes. Through this molecular interaction, Trim71 cooperates with ESC-specific miR-290 and miR-302 to bind to the 3’UTR of Cdkn1a and inhibit its translation. Because Cdkn1a is a negative regulator of the G1-S transition, the miRNA-dependent 3’UTR activity of Trim71 promotes the cell cycle structure necessary for ESC self-renewal. Importantly, Trim71 function is tightly incorporated into a larger intracellular network because it is a target of the prodifferentiation miRNA let-7, whose biogenesis is inhibited by the RBP Lin28a (see the subsection “Multifunctional RBPs,” below). Adding to the complexity of this regulatory circuit, another study found that Trim71 is also an E3 ubiquitin ligase for Ago2 and showed through luciferase assays that Trim71 cooperates with Lin28a to inhibit let-7 activity (Rybak et al., 2014). Using deletion mutants, both of these groups clearly establish Trim71 as a key player for ESC self-renewal. Pum1 disruption promoted self-renewal in an ESC-like state when the cells were cultured in conditions permissive to differentiation (Leeb et al., 2014).

Another RBP that destabilizes pluripotency transcripts is the AU-rich element mRNA-binding protein (AUBP) Brf1. AUBPs are known to be developmentally important regulators of splicing, stability, translation, and localization, and Brf1 is critical in embryogenesis: knockout mice die by E10.5 (Stumpo et al., 2004). A recent study found that activation of the FGF-Erk signaling pathway, which stimulates mouse ESC differentiation, leads to upregulation of Brf1 (Tan and Elowitz, 2014). Brf1 overexpression under pluripotency conditions impaired ESC proliferation, and under differentiation conditions it promoted the acquisition of mesendoderm fate. RNA immunoprecipitation and sequencing (RIP-seq) showed that Brf1 binds to AU-rich elements (AREs) in a number of pluripotency miRNAs, including Nanog, Klf2, Kdm4c, and Zfp143. Moreover, FGF stimulation led to degradation of Nanog transcripts in an ARE-dependent fashion, implicating Brf1 in miRNA destabilization and ESC differentiation. Future work will likely reveal more roles for AREs and other AUBPs in pluripotency and early embryonic development.

With all of these RBPs involved in determining the stability of different mRNA transcripts, it will be interesting to explore to what extent their targets overlap and whether they cooperate to coregulate a particular subset of transcripts. For those mRNAs with binding sites for several RBPs, it would furthermore be worthwhile to examine how the relative RBP concentrations and motif binding affinities affect transcript levels.

**Regulation of Translation**

RBPs can also directly modulate protein translation, often through binding at the 5’UTR of RNA transcripts. In so doing, RBPs recruit translation initiation factors, adjust the accessibility of the RNA to ribosomes, create ribonucleoprotein structures conducive for cap-independent and internal ribosome entry site (IRES)-mediated translation, and regulate the movement of the ribosomes along the transcript. Given these many roles, it is no surprise that RBPs involved in these 5’UTR-related processes have also been linked to the control of ESC pluripotency. In particular, RBP-5’UTR interactions involving the proteins Nat1, Rbm35a (Esrp1), and Ptbp1 have been shown to regulate ESC differentiation and proliferation.

Nat1 was first identified in liver carcinomas as a general repressor of translation. In mouse ESCs, depletion of Nat1 does not affect proliferation and self-renewal in ESC growth conditions, but Nat1−/− ESCs exhibit a defect in retinoic acid (RA)-induced differentiation (Yamanaka et al., 2000). Previous studies in other cell lines have shown that Nat1 is homologous to the translation initiation factor eIF4G, is located in the cytoplasm and plays a crucial role in maintaining the translation machinery.
and can autoregulate its own translation from an IRES in its mRNA; thus, the authors speculated that Nat1 binds to highly structured 5′UTRs and associates with translational initiation factors. However, their study did not detect a difference in cap-dependent or cap-independent translation in wild-type and Nat1+/− ESCs as measured by [35S]methionine incorporation and bicistronic luciferase assays, respectively. Despite the negative results, it is important to note that these experiments were conducted under steady-state pluripotency conditions, not under the differentiation conditions in which they observed the phenotype. Therefore, it remains plausible that Nat1 influences ESC differentiation through translational mechanisms, and further studies should address the role of Nat1 in transitions into and out of pluripotency.

Like Nat1, Rbm35a is a negative regulator of pluripotency. Knockdown of Rbm35a inhibits ESC differentiation and promotes somatic cell reprogramming by increasing expression of Oct4, Nanog, and Sox2. Indeed, Rbm35a immunoprecipitation and polysome profiling show that Rbm35a normally binds to the 5′UTR of Oct4 and Sox2 mRNAs, thus preventing them from being loaded into polysomes (Fagoonee et al., 2013). In addition, Rbm35a activity is not restricted to ESCs, because the RBP is a regulator of alternative splicing in epithelial cell lines (Warzeca et al., 2009) and has also been shown to be expressed in tumor cells where it binds to highly structured GC-rich 5′UTRs (Warzecha et al., 2009) and has also been shown to be expressed in tumor cells where it binds to highly structured GC-rich 5′UTRs of oncogenes and prevents their translation (Leonieva and Ionov, 2009). ESCs themselves form teratomas when injected into mice and share many properties with somatic tumors, including limitless replicative potential. Therefore, Rbm35a suppression may be an important component of maintaining ESC and tumor cell immortality through both shared and unique molecular targets present in these different cellular contexts.

Ptbp1, another RBP that binds the 5′UTR of gene transcripts, also appears to control ESC growth through its regulation of the cell cycle. Ptbp1 knockout ESCs have a proliferation defect with a prolonged G2/M phase. This phenotype appears to be at least in part secondary to problems with chromosomal segregation. Bicistronic luciferase assays show that Ptbp1 binds to the IRES of CDK11P58 and represses translation of this gene, high levels of which are associated with a prolonged telophase caused by chromosomal lagging. Overexpression of CDK11P58 in wild-type ESCs led to chromosome missegregation (Ohno et al., 2011). Nevertheless, it is unclear whether or not overexpression of CDK11P58 is the sole contributor to the Ptbp1 knockout phenotype—a question that could be addressed through global comparisons of mRNA and protein abundance in wild-type and knockout cells. Notably, Ptbp1 has also been extensively studied in other cell lines, where it has been implicated in alternative splicing, alternative polyadenylation, mRNA stability at the 3′UTR, and IRES-driven translation (Boultz et al., 2007; Castelo-Branco et al., 2004; Kosinski et al., 2003; Bushell et al., 2006). It would be useful and informative for future studies to examine whether these other well-known functions of Ptbp1 are also involved in pluripotency.

Multifunctional RBPs

In addition to Adar, Pum1, Rbm35a, and Ptbp1, other RBPs have been proposed to play pleiotropic roles. For example, while Lin28a clearly regulates let-7 biogenesis through directing uridylation of pre-let-7, it appears to have additional functions independent of its relationship with let-7 (reviewed in Shyh-Chang and Daley, 2013). Through binding to an AG-rich motif similar to its pre-let-7 binding motif (Wilbert et al., 2012; Cho et al., 2012), Lin28a both positively and negatively modulates the translation of hundreds of mRNAs in human and mouse ESCs (Wilbert et al., 2012; Cho et al., 2012; Peng et al., 2011; Xu et al., 2009). How Lin28a achieves this incredibly complex orchestration of its many targets remains to be determined.

Lin28a is an unusual example of an RBP attributed with multiple functions in a single cellular context. To date, most studies done on multifunctional RBPs, like Ptbp1 and Adar, only investigate a single function of that RBP in ESCs, which does not exclude the possibility that the RBP’s other proposed functions are also important in supporting the pluripotency network. In general, a number of RBPs, such IIf2, Unr, and others, have been genetically deleted in ESCs and shown to have an effect on the ESC phenotype, but currently, the functions of these RBPs can only be inferred from work done in other cell types. As every study ascribes a different role to each of these RBPs, it is unclear which, if any, are actually involved specifically in the context of pluripotency.

A notable example providing evidence that RBPs can be multifunctional within a single context comes from a recent study of the RNA helicases DDX5 and DDX17 not in pluripotent cells but in the differentiation of myoblast and epithelial cell lines (Dardenne et al., 2014). The authors showed that DDX5 and DDX17 cooperate with the hnRNP H/F splicing factors to promote a differentiation-specific splice profile. At the same time, DDX5 and DDX17 serve as transcriptional coregulators of key differentiation transcription factors as well as miRNAs that, in turn, negatively feed back on the expression of the RBPs themselves. It is unclear whether these two roles for DDX5 and DDX17 are physically linked; nevertheless, this story provides evidence that an RBP can simultaneously function at the level of DNA (transcription) and RNA (splicing).

Conclusions and Perspectives

It is without a doubt that posttranscriptional regulation by RBPs contributes extensively to the establishment and maintenance of, as well as exit from, the ESC state. The field remains relatively young and largely uncharted, however, and there are many opportunities for further inquiry and discovery.

As described above and summarized in Table 1, a number of studies have documented the significance of individual RBPs in pluripotency based on knockdown and knockout models demonstrating that perturbation of the RBP disrupts the wild-type ESC phenotype. However, few probe the actual mechanism by which the RBPs produce their effect in ESCs. Instead, RBP function has been examined mostly in cancer cell lines and cell-free biochemical assays, the in vivo relevance of which needs to be more clearly characterized. The studies that do delve into the molecular mechanisms in ESCs generally examine only a limited number of targets. While the RBPs discussed here are not unique to ESCs, cell context likely influences their downstream effects on cell fate and cell fate transitions significantly by providing a specific combination of intracellular pathways with which these RBPs interface. Thus, to reach a deep and unified understanding of the role of RBPs
in pluripotency, we will need to make use of global approaches like those that have already been successfully applied to the field of epigenetics. In particular, we must dissect all RBP-RNA interactions systematically in ESC self-renewal and differentiation conditions so as to examine not only how RNA transcript levels change but also how the components of

| Table 1. Summary of RBPs Associated with the Pluripotency Network |
|---------------------------------|--------|---------------------------------|---------------------------------|
| Category of Posttranscriptional Regulation | RBP | Positive or Negative Regulator of Pluripotency? | Mechanism of Action in ESCs | Other Functions (Not Validated in ESCs) |
| Alternative splicing | FOX2 (RBM9, FXH) | positive (Yeo et al., 2009) | binds upstream of splice site for exon exclusion; binds downstream of splice site for exon inclusion | |
| | SON | positive (Lu et al., 2013) | binds short introns with weak splice sites | |
| | MBNL1, MBNL2 | negative (Han et al., 2013) | binds upstream of splice site for exon inclusion; binds downstream of splice site for exon exclusion | |
| | SFRS2 (SC35) | positive (Lu et al., 2014) | regulates splicing of MBD2 | |
| Alternative Polyadenylation | Fip1 | positive (Lackford et al., 2014) | promotes ESC-specific patterns of alternative polyadenylation | |
| RNA modification | Zcchc11, Zcchc6 (TUTases) | positive (Hagan et al., 2009) | inhibits let-7 biogenesis by uridylylating the pre-miRNA | mRNA stability with Elav1 (HuR); A-to-I editing of pri-miRNAs to prevent their processing; promote pre-miRNA cleavage in complex with Dicer |
| | ADAR (ADAR1) | positive (Osenberg et al., 2010) | drives A-to-I editing | |
| | Mettl3, Mettl14 | positive (Wang et al., 2014a) | methylates adenosines in mRNA, reducing transcript stability | |
| Nuclear export | Thoc2, Thoc5 (members of THO complex) | positive (Wang et al., 2013b) | exports pluripotency gene transcripts to the cytoplasm for translation | |
| RNA stability | Trim71 (mLin41) | positive (Chang et al., 2012; Rybak et al., 2009; Loedige et al., 2013; Worringer et al., 2014) | binds Ago2 complexes, enhancing ESC miRNAs’ inhibition of Cdkn1a expression, which thus promotes ESC cell cycle structure; antagonizes Ago2 by ubiquitination; degrades prodifferentiation gene transcripts | |
| | Pum1 | negative (Leeb et al., 2014) | degrades pluripotency gene transcripts | translation repressor |
| | Brf1 | negative (Tan and Elowitz, 2014) | degrades pluripotency gene transcripts | |
| Translation | Nat1 | negative (Yamanaka et al., 2000) | unclear | translation repressor |
| | Rbm35a | negative (Fagoonee et al., 2013) | prevents polysome loading of pluripotency gene transcripts | alternative splicing |
| | Ptbp1 | positive (Ohno et al., 2011) | binds to IRES and represses translation of CDK11p58 | alternative splicing; alternative polyadenylation; mRNA stability at 3’UTR |
| Multifunctional | Lin28a | positive (reviewed in Shyh-Chang and Daley, 2013) | inhibits let-7 biogenesis by recruiting TUTases to uridylylate pre-let-7; translational activation; mRNA degradation with Drosha | |
Intriguingly, as mentioned above under the subsection “Multi-functional RBPs,” a number of RBPs appear to be involved in multiple aspects of RNA metabolism in both the nucleus and the cytoplasm. This observation may result from an RBP having different roles in different cell types, multiple roles in a single cell type, or some mix of the two. Regardless, it is almost certain that the particular combination of targets and cofactors that an RBP encounters influences its functions in context-specific ways—a notion that expands upon the “RNA regulon” model originally proposed by Jack Keene, in which an RBP binds multiple targets to effect changes in various cellular processes (Keene, 2007). In other words, it is possible that instead of regulating multiple targets through just a single mechanism, one RBP could simultaneously participate in several layers of RNA metabolism (Figure 2). In so doing, the RBP becomes part of increasingly complex regulatory modules, with many opportunities for feedback and crosstalk, where one aspect of metabolism such as translation can be linked to another such as splicing. Consequently, perturbation of the RBP could have a cascading effect on the molecular landscape of a cell and precipitate drastic switches in cellular identity. Also, it follows that disrupting subsets of the mRNA targets or cofactors of any one RBP could affect parallel pathways by shifting the RBP’s dominant activities to different genes or even different regulatory modules altogether. For example, a decrease in the levels of a cofactor that enables an RBP to regulate splicing could drive that RBP to shift its main activity to transcription.

Moving forward, it will be critical to investigate RBPs at the genomics level in biologically relevant cell types, focusing on those states in which the RBPs are demonstrated to play significant roles. Studying RBPs in ESCs provides an excellent starting context to achieve such a goal, because not only can ESCs provide unlimited untransformed material for large-scale genomic studies, they can also be differentiated down any cellular lineage to determine how context changes protein function. Thus, experimental tools and platforms developed in the ESC system can be used to study a multitude of cell types that comprise the mammalian body plan.

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