

Series: Fresh Perspectives from Emerging Experts

# Review

# Linking Long Noncoding RNA Localization and Function

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Recent studies have revealed the regulatory potential of many long noncoding RNAs (IncRNAs). Most IncRNAs, like mRNAs, are transcribed by RNA polymerase II and are capped, polyadenylated, and spliced. However, the subcellular fates of IncRNAs are distinct and the mechanisms of action are diverse. Investigating the mechanisms that determine the subcellular fate of IncRNAs has the potential to provide new insights into their biogenesis and specialized functions.

#### The Diversity of Long Noncoding RNAs

By their broadest definition, long noncoding RNAs (IncRNAs) comprise different types of RNA polymerase II (Pol II)-transcribed molecules with sizes greater than 200 nt in length, which distinguishes them from short RNAs such as microRNAs (Figure 1). It is now widely accepted that IncRNAs have regulatory roles in gene expression at both the transcriptional and posttranscriptional levels in diverse cellular contexts and biological processes. IncRNAs are responsible for nuclear structure integrity, and can regulate the expression of either nearby genes (acting in cis in the nucleus) or genes elsewhere in cells (acting in trans in the nucleus or cytoplasm) by interacting with proteins, RNAs, and DNAs. IncRNAs can positively or negatively regulate gene expression through multiple mechanisms, including through recruiting transcription factors or chromatin-modifying complexes to their DNA targets, by forming heterogeneous nuclear ribonucleoprotein (hnRNP) complexes, by acting as decoys to sequester RNA-binding proteins (RBPs) and microRNAs, or by directly interacting with RNAs and DNAs by base pairing (see reviews [1-3]).

Most annotated IncRNAs are Pol II transcribed, hence they are presumably capped, polyadenylated, and spliced just like mRNAs [4,5]. Such IncRNAs are transcribed either from intergenic regions of the genome [i.e., large intergenic noncoding RNAs (lincRNAs); Figure 1A] or from the opposite strand of protein-coding genes [natural antisense transcripts (NATs); Figure 1B] [6]. lincRNAs are the most abundant class of IncRNAs comprising over 10 000 species so far. A major difference between these IncRNAs and mRNAs is that IncRNAs have little if any proteincoding potential. IncRNAs may contain fewer exons than mRNAs and often have weak cryptic splicing and polyadenylation signals [5].

While a large proportion of IncRNAs look like mRNA transcripts, a number of IncRNAs form their 3' end in unusual ways (Box 1 and Figures 1C-F) and others are processed from long polyadenylated primary transcripts (Box 1 and Figures 1G-I). In addition, new types of IncRNAs are likely to be discovered through integrated approaches. Recently, the advent of RNA-centric methods combined with deep sequencing as well as emerging genome-editing technologies have greatly advanced our knowledge of both the functional significance and mechanisms of action of IncRNAs (see reviews [3,7]). Despite these rapid areas of growth, we still know little about the life cycle of most 'mRNA-like' IncRNAs. How their fates and functions are different from each other and how cells distinguish IncRNAs from mRNAs remain obscure. Importantly, it is

#### **Trends**

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The function of IncRNAs is associated with their unique subcellular localization

Mature IncRNAs can accumulate in cis, localize in the nucleus in trans. or export to the cytoplasm to execute their functions.

Multiple factors including ribonucleic nuclear retention elements nuclear protein factors, higher-order chromosome organization, and the coupling of RNA-protein assemblies with IncRNA transcription may dictate the subcellular localization patterns of IncRNAs.

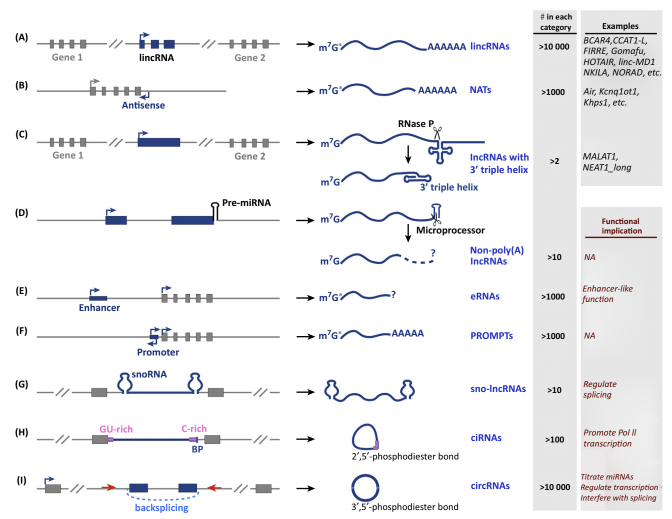
A better understanding of the IncRNAs themselves is crucial to link these noncoding transcripts to RNA biology and to address their cellular roles in depth.

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Figure 1. The Diversity of Long Noncoding RNAs (IncRNAs) in Mammalian Genomes. (A) Large intergenic noncoding RNAs (lincRNAs). (B) Natural antisense transcripts (NATs). (C and D) IncRNAs with alternatively processed 3' ends by ribonuclease P (RNase P; C) or microprocessor (D) cleavage. (E) Enhancer RNAs (eRNAs). (F) Promoter upstream transcripts (PROMPTs). (G) Small nucleolar RNA (snoRNA)-ended IncRNAs (sno-IncRNAs). (H) Circular intronic RNAs (ciRNAs). ciRNA formation depends on consensus RNA sequences (pink bars) to avoid debranching of the lariat intron. (I) Circular RNAs (circRNAs) produced from backsplicing of exons. Inverted complementary sequences (red arrows) in introns flanking circularized exons promote circRNA biogenesis. Gray columns: left, the number of lncRNAs identified in each category in mammals so far; right, examples and functional implication of lncRNAs of each category. \*These Pol II transcripts are presumably capped, but the direct lines of evidence showing the presence of m<sup>7</sup>G cap at their 5' termini are still warranted. Abbreviations: BCAR4, Breast cancer antiestrogen resistance 4; BP, branch point; CCAT1-L, colon cancer associated transcript 1-long isoform; FIRRE, functional intergenic repeating RNA element; HOTAIR, HOX transcript antisense RNA; NKILA, nuclear factor-κB interacting IncRNA; NORAD, noncoding RNA activated by DNA damage; Pol II, polymerase II.

becoming increasingly clear that the function of IncRNAs is associated with their unique subcellular localization, but how cells sort different IncRNAs to specific subcellular compartments for carrying out their roles in gene regulation remains unclear. Understanding the mechanisms of how the subcellular fate of IncRNAs is achieved and regulated has the potential to provide new insights into their biogenesis and functions.

#### The Function of IncRNAs Are Associated with Their Subcellular Fates

Like proteins, the function of IncRNAs depends on their subcellular localization. Many IncRNAs are recognized as important modulators for nuclear functions (see reviews [8,9]) and exhibit distinct nuclear localization patterns (Figures 2A-D). Others must be exported to the cytoplasm



#### Box 1. Processing of Long Noncoding RNAs (IncRNAs) in Alternative Ways.

First, MALAT1 and NEAT1\_long are two IncRNAs that are localized to nuclear speckles [68] and paraspeckles [31], respectively. They are processed at their 3' ends by recognition and cleavage of tRNA-like structures by ribonuclease P (which processes the 5' ends of tRNAs). Ribonuclease P cleavage leads to the formation of their mature 3' ends, which are subsequently protected by a conserved stable U-A•U triple-helical RNA structure ('•' denotes the Hoogsteen face and '-' denotes the Watson-Crick face) [60,61] (see Figure 1C in main text). Second, some IncRNA transcripts containing miRNAs (Inc-pri-miRNAs) use microprocessor cleavage to terminate transcription, resulting in unstable IncRNAs without 3' end poly(A) tails [69] (see Figure 1D in main text). Third, enhancer RNAs (eRNAs) are bidirectionally transcribed from enhancers by polymerase II (Pol II) and lack poly(A) tails (see Figure 1E in main text) [70,71] but have enhancer-like functions [72-74]. Depletion of Integrator, a complex with an RNA endonuclease activity, resulted in the accumulation of primary eRNA transcripts that remained bound to Pol II, suggesting a role of Integrator in cleaving the 3' end of eRNAs [75]. Fourth, promoter upstream transcripts (PROMPTs) are bidirectionally transcribed from promoters of protein-coding genes. They are 200-600 nt in length, capped, and polyadenylated (see Figure 1F in main text). They are nuclear retained and undergo rapid degradation [76,77]. PROMPTs may not have a function. However, the rapid degradation of transcribed PROMPTs may relate to the choice of promoter directionality [78]. Both PROMPTs and eRNAs appear to be degraded by the RNA exosome [79]. Fifth, excised introns can also generate IncRNAs. Small nucleolar RNA-ended IncRNAs (sno-IncRNAs) are formed when one intron contains two snoRNA genes (see Figure 1G in main text). During splicing, the sequences between snoRNAs are not degraded, leading to the accumulation of IncRNAs flanked by snoRNAs but lacking 5' caps and 3' poly(A) tails in the nucleus [20]. Their expression is species specific and results from species-specific alternative splicing [80]. Finally, circular RNAs are yet another type of IncRNAs that are protected from RNA exonucleolytic decay machineries. They can be derived from the inefficient debranching of lariat introns (see Figure 1H in main text) [63] or from pre-mRNAs backsplicing of exons (see Figure 1I in main text). Circular RNAs may regulate gene expression by titrating miRNAs, regulating transcription, or interfering with splicing (see review [81]).

to carry out their regulatory roles (Figure 2E). In this review, a number of well-characterized IncRNAs are classified into three groups depending on their subcellular localization to illustrate the association of IncRNA localization and function: those that are absolutely nuclear localized in cis (Figures 2A,B), those that are mainly nuclear localized and function in trans (Figures 2C,D), and those that largely localize and function in the cytoplasm (Figure 2E). It is worth noting that a recent large-scale evaluation of the subcellular fates of IncRNAs in human cell lines using singlemolecule RNA fluorescence in situ hybridization revealed that IncRNAs exhibited a wide range of subcellular localization patterns, including not only distinct patterns of nuclear localization but also nonspecific location in both the nucleus and cytoplasm [10].

#### IncRNAs That Accumulate in cis

Of those that have nuclear functions, one group of IncRNAs accumulates at their sites of transcription (in cis; Figures 2A,B). These IncRNAs can be classified into two subgroups based on whether they act in cis, close to the site of transcription, or in trans. The former IncRNAs execute their regulatory roles in cis by various mechanisms (Figure 2A). For example, they can modulate local gene expression either by recruiting transcription factors, chromatin organizers, or chromatin modifiers, or by forming a DNA-RNA triplex that anchors the IncRNA and associated effector proteins to the gene promoter. For example, a set of IncRNAs is induced by DNA damage from the promoter of cyclin D1 gene (CCND1); these IncRNAs bind to the translocated in liposarcoma (TLS) protein and allosterically modulate TLS activity to inhibit histone acetyltransferase in cis, which suppresses CCND1 transcription [11]. Furthermore, regulatory RNAs transcribed from promoter-proximal and promoter-distal enhancers remain bound to chromatin and contribute to stable transcription factor (such as YY1) occupancy at these sites to maintain geneexpression programs [12]. In addition, the IncRNA CCAT1-L is transcribed from the upstream super-enhancer region of the MYC gene; CCAT1-L accumulates in cis, interacts with the chromatin organizer CTCF, and promotes chromosome looping between the super-enhancer and the MYC promoter, resulting in enhanced MYC transcription in colorectal cancer [13].

Several well-characterized NATs accumulate to their sites of transcription and affect expression of neighboring genes. In mammalian imprinting, Air and kcng1ot1 IncRNAs are involved in the allelespecific silencing of their cis-linked genes [14-16]. Air recruits H3K9 histone methyltransferase G9a to maintain the imprinted silencing of multiple genes including the lgf2r locus [14].



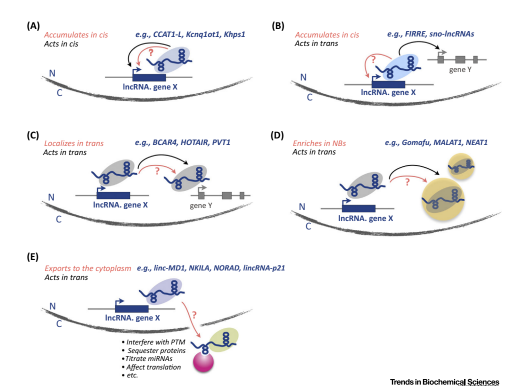


Figure 2. Functions of Long Noncoding RNAs (IncRNAs) Are Associated with Their Subcellular Fates. IncRNAs have distinct subcellular localization patterns, allowing IncRNAs to execute their specified functions. (A) IncRNAs can accumulate and act in cis once they are transcribed. (B) IncRNAs can accumulate in cis once they are transcribed, but act in trans that affects genes located in the same chromosome at a distance or in different chromosomes. (A and B) Mechanisms of IncRNA in cis localization are largely unknown. (C) IncRNAs can localize to elsewhere in the nucleoplasm in trans and act in trans. (D) IncRNAs can accumulate to specific nuclear bodies (orange circles) and act in trans. The mechanisms of IncRNA nuclear retention remain largely unknown, as does whether such IncRNAs are required to be translocated in the nucleus to regulate gene expression. (E) IncRNAs can be exported to the cytoplasm to execute their functions. For example, a cytoplasmic IncRNA can sequester protein (pink circle) or interfere with protein post-translational modification (PTM). Whether the nucleocytoplasmic export of cytoplasmic IncRNAs is distinct from that of mRNA has not yet been examined. The color of shaded oval indicates differences in protein composition of long noncoding ribonucleoproteins. Pink arrows, the unknown mechanisms for IncRNAs to gain specific subcellular localization patterns; black arrows, IncRNAs execute functions in distinct subcellular compartments. BCAR4, breast cancer antiestrogen resistance 4; CCAT1-L, colon cancer associated transcript 1-long isoform; FIRRE, functional intergenic repeating RNA element; lincRNA, large intergenic noncoding RNAs; MALAT1, metastasis-associated lung adenocarcinoma transcript 1; NBs, nuclear bodies; NEAT1, nuclear-enriched abundant transcript 1; NKILA, nuclear factor-κΒ interacting IncRNA; NORAD, noncoding RNA activated by DNA damage; PVT1, plasmacytoma variant translocation 1; Sno-IncRNA, small nucleolar RNA-ended IncRNAs.

By contrast, kcnq1ot1 recruits G9a and the H3K27 histone methyltransferase PRC2 (polycomb repressive complex 2), which leads to the bidirectional silencing of a dozen paternally imprinted genes in the Kcnq1 locus [15,16]. Finally, the antisense IncRNA Khps1 activates proto-oncogene SPHK transcription in an E2F1-dependent manner. This is achieved by forming a DNA-RNA triplex of the SPHK1 promoter with Khps1, which further recruits the histone acetyltransferase p300/CBP to the SPHK1 promoter, leading to changes in chromatin structure and enhancing E2F1-dependent expression of SPHK1 [17]. Importantly, the act of transcription itself may influence adjacent gene expression in addition to the effects of the in-cis-accumulated IncRNA products. For instance, the imprinted Igf2r silencing also requires Air transcriptional overlap of the Igf2r promoter, which interferes with Pol II recruitment in the absence of repressive chromatin [18].

In-cis-localized IncRNAs can also influence gene expression in trans (Figure 2B). The IncRNA FIRRE is transcribed from the X chromosome and involved in the formation of five transchromosomal contacts at its site of transcription [19]. This spatial colocalization is achieved



through the 3D organization of the chromosomes, which enables long-range interactions between the regulatory genomic elements and the target gene locus. Such colocalization patterns are dramatically decreased in cells that lack the Firre locus [19]. In addition, there is yet another mechanism through which in-cis-accumulated IncRNAs can act as decoys for RBPs to achieve gene regulation in trans. For example, in human embryonic stem cells, small nucleolar RNA-ended IncRNAs (sno-IncRNAs) expressed from chr15q11-q13 are abundant and accumulate near their sites of synthesis. They sequester Fox2 and might influence Fox2-mediated alternative splicing regulation in trans [20].

The mechanism through which these IncRNAs are restricted to their sites of transcription is unknown. Binding to local factors [11-16] or forming RNA-DNA triplex structures [17] may facilitate IncRNAs accumulation in cis. Another attractive hypothesis that remains to be tested is that these IncRNAs may remain bound to Pol II [12] and their in cis localization is somehow coupled to Pol II transcription.

#### IncRNAs That Localize in the Nucleus in trans

A second group of IncRNAs also localizes to the nucleus, but they often need to be relocated from their sites of synthesis to have an impact on gene regulation, which can be either globally or in a gene-specific manner (Figure 2C). HOTAIR is one of the best-studied examples and can be used to illustrate this group of IncRNAs. HOTAIR is transcribed from the mammalian HOX C locus, but it represses transcription in trans at the HOX D locus by recruiting PRC2 during developmental patterning [21,22]. The expression of HOTAIR also affects the cancer epigenome by genome-wide retargeting of PRC2, leading to altered histone H3 lysine 27 trimethylation (H3K27me3), a chromatin mark that is indicative of transcription repression [23]. In addition to the HOX D locus, HOTAIR binds to hundreds of other genomic sites through a GA-rich motif to nucleate broad domains of polycomb occupancy and H3K27me3 [24]. The GA-rich HOTAIR motif may enable formation of an RNA:DNA triplex that recruits the HOTAIR-chromatin modification complex. BCAR4 and PVT1 are two other IncRNAs that predominately localize in the nucleoplasm. BCAR4 binds to transcription factors SNIP1 and PNUTS via its two distinct regions and alters downstream epigenetic regulation in response to chemokine signals in breast cancer [25]. PVT1 interferes with phosphorylation of MYC at Thr58 in the nucleus, which subsequently increases MYC stability and leads to elevated MYC levels in cancers [26].

Whereas HOTAIR, BCAR4, and PVT1 have a global localization within the nucleus, some other nuclear IncRNAs accumulate within specific nuclear bodies (Figure 2D). These membrane-less subnuclear structures contain specific protein and IncRNA components that define particular nuclear processes. Gomafu (also referred to as MIAT in humans) localizes to distinct nuclear domains in specific neuronal cells and associates with RBPs including Celf3 and SF1 [27], suggesting of role of Gomafu in sequestering proteins. MALAT1 localizes to nuclear speckles, which contain various splicing factors [28], and has been recently shown to regulate mammary cancer pathogenesis [29]. The NEAT1 IncRNAs, expressed from human chr.11q13.1, are abundant IncRNAs that are localized to paraspeckles [30]. There are approximately 5-20 paraspeckles in each nucleus of most mammalian cells and NEAT1 is required for the integrity of paraspeckles [30-33]. NEAT1 interacts with multiple paraspeckle-localized proteins [34,35] and hundreds of genomic sites of active genes in human cells [35]. Although how NEAT1 and paraspeckles are assembled and localized has yet to be defined precisely, it is clear that paraspeckle assembly occurs during the transcription of NEAT1 [36] and that a change in the transcription of NEAT1 leads to altered paraspeckle formation [37]. Hence, coupling with Pol Il transcription to nucleate nuclear domains represents one way that IncRNAs can nonrandomly accumulate in the nucleus. How other IncRNAs accumulate in trans is largely unknown. Since roles of this group of IncRNAs are diverse, mechanisms of their nuclear localization are expected to vary.



#### IncRNAs That Are Exported to the Cytoplasm

The third group of IncRNAs needs to be exported to the cytoplasm to carry out their regulatory roles (Figure 2E). Increasing lines of evidence show that such IncRNAs can affect gene expression in multiple ways. Cytoplasmic IncRNAs can interfere with protein post-translational modifications, resulting in aberrant signal transduction [38,39]. For example, the IncRNA NKILA interacts with NF-κB and interferes with phosphorylation of IκB. This leads to inhibition of IKKinduced IκB phosphorylation and NF-κB activation and suppression of breast cancer metastasis [38]. Furthermore, cytoplasmic IncRNAs can influence gene regulation by acting as decoys for miRNAs [40] and proteins [41]. It has been shown that linc-MD1 sponges miR-133, which usually targets transcription factors to activate muscle-specific gene expression [40]. A highly conserved and abundant IncRNA NORAD regulates genomic stability by sequestering PUMILIO proteins, which represses the stability and translation of mRNAs to which they bind [41]. Moreover, other IncRNAs can also affect mRNA translation in the cytoplasm [42,43]. STAU1-mediated mRNA decay (SMD) involves the degradation of translationally active mRNAs whose 3'-untranslated regions (3'-UTRs) bind to STAU1, a protein that binds to doublestranded RNA (dsRNA). The requisite dsRNA can be formed by cytoplasmic IncRNAs base pairing with a complementary Alu embedded within the mRNA 3'-UTRs [42]. lincRNA-p21 also has a role in mRNA translation suppression. In the cytoplasm, HuR influences lincRNA-p21 decay; in the absence of HuR, lincRNA-p21 is stable and identifies mRNA targets including JUNB and CTNNB1 mRNAs by base pairing and represses their translation [43].

Finally, recent proteomic analyses revealed that a subset of transcripts currently annotated as IncRNAs may nevertheless encode small proteins (see review [44]) and several annotated IncRNAs can indeed encode functional micropeptides [45-47]. Thus, functional studies of cytoplasmic IncRNAs should also carefully assess whether the IncRNA of interest has any coding potential. In this scenario, one advantage of studying nuclear IncRNAs is that they are excluded from the possibility of being translated, therefore assuring confidence that the experimental conclusions reached truly reflect RNA functions.

Together, these emerging studies reveal that it is critical for IncRNAs to be localized to specific subcellular compartments for their functions. In the next section, the mechanisms that may account for the distinct subcellular fates of different IncRNAs are discussed, focusing on nuclearretained ones. This area contains many aspects that have not been well addressed in metazoans.

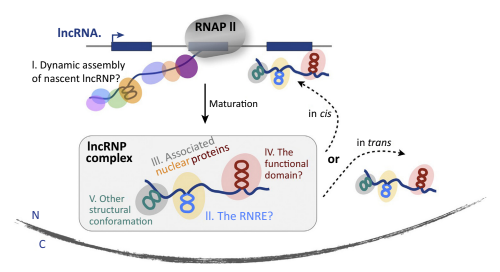
#### Mechanisms of IncRNA Subcellular Fate Determination

At the chromatin level, IncRNA loci appear to follow the same rules as for protein-coding genes, although there are some differences between chromatin assemblies at divergent promoters for sense mRNA versus antisense IncRNAs in yeast [48]. These similarities led to the systematic discovery of IncRNAs by monitoring the patterns of histone 3 Lys 4 trimethylation (H3K4me3) in the promoter region followed by histone 3 Lys 36 trimethylation (H3K36me3) across the actively transcribed intergenic regions in mammalian cells. The same patterns are also associated with protein-coding genes [3,4,49,50]. These studies indicate that the chromatin status is unlikely to be a major factor in either distinguishing the fate and function of IncRNAs from mRNAs or determining the diverse cellular localizations among IncRNAs. Other than chromatin-mediated regulation, several possible mechanisms may dictate the subcellular localization patterns of IncRNAs.

### Nuclear Retention Signals in IncRNAs Are Deposited during Transcription through the Assembly of Long Noncoding Ribonucleoproteins Complexes

An interesting observation is that IncRNAs produced by alternative processing pathways (Box 1 and Figures 1C-G) are likely to be enriched in the nucleus in mammalian cells, suggesting that the processing of IncRNAs may be associated with their distinct subcellular localization and





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Figure 3. Mechanisms of Long Noncoding RNA (IncRNA) Nuclear Retention. Multiple factors can be involved in IncRNA nuclear retention. The nuclear retention signals of IncRNAs might be deposited during the assembly of nascent long noncoding ribonucleoproteins (IncRNPs) complexes (I). Upon IncRNP maturation, ribonucleic nuclear retention elements (RNREs; II) residing in IncRNAs may associate with nuclear matrix proteins (III, yellow oval) to constrain IncRNAs in the nucleus. In addition, functional domains (IV) of IncRNAs may recruit chromatin-modifying complexes or transcription factors, or other structural conformation of IncRNAs (V) that can subsequently keep these IncRNA molecules from export to the cytoplasm and facilitate their nuclear localization in cis or in trans. Abbreviation: RNAP, RNA polymerase II.

function, for instance, the lack of poly(A) tail may inhibit IncRNA export. However, whether the transcription and processing of mRNA-like IncRNAs (Figures 1A,B) are linked to their distinct subcellular localization and function is awaiting exploration.

Pol II transcription, nascent pre-mRNA splicing, capping, polyadenylation, mRNA export, and surveillance are seamlessly integrated during mRNA maturation. Throughout this process, premRNAs are decorated with various proteins to form higher-order messenger ribonucleoprotein (mRNP) complexes. Mature mRNPs are then exported to the cytoplasm where translation and mRNA turnover take place (see review [51]).

To distinguish IncRNAs from nuclear-retained mRNAs or cytoplasmic IncRNAs, nuclear retention signals may decorate the nuclear-retained IncRNAs upon transcription. In this model, the sequential assembly of the long noncoding ribonucleoprotein (IncRNP) complexes of future nuclear-retained IncRNAs during their transcription would be substantially distinct from that of mRNPs or that of IncRNPs that needed to be exported to the cytoplasm (Figure 3). Some components of IncRNPs are likely to be different from those of mRNPs. Such dynamic assembly of IncRNPs during transcription and processing may in turn direct their specific subnuclear localization patterns.

Although there is no established evidence yet to support this model in mammalian cells, recent studies in yeast began to reveal differences between the biogenesis of IncRNAs and mRNAs. In the budding yeast Saccharomyces cerevisiae, the transcription preinitiation complexes and key transcription factors that regulate IncRNA and mRNA expression appear to have no detectable differences [52]. Rather, later steps in transcription appear to provide a divergence point. By comparing the maturation pathways of mRNAs and different types of IncRNAs, such as stable unannotated transcripts (SUTs) and cryptic unstable transcripts (CUTs), in budding yeast, Tuck and Tollervey [53] observed that the cellular fate of IncRNA and mRNA transcripts is largely



determined during 3' end formation prior to the acquisition of export competence [53]. The nuclear RNA surveillance machinery rapidly targets CUTs terminated by Nrd1-dependent transcription termination, whereas SUTs contain signals for cleavage and polyadenylation just like mRNAs. Moreover, while most SUTs remain in the nucleus, a subset can be exported to the cytoplasm [53]. However, it remains to be determined whether similar features can be applied to IncRNAs in higher eukaryotic cells. As mRNP transcription, processing, and export are much more complex in mammals, it is likely that the regulation of lncRNA production during transcription and processing in these cells is more complicated than is currently appreciated in yeast. For example, the CCAT1 locus produces two abundant lncRNAs with overlapping 5' ends: CCAT1-L (the 4700-nt long isoform) and CCAT1-short isoform (CCAT1-S; the 2600-nt short isoform) that is proposed to be generated by 3' end processing of the long CCAT1 in colorectal cancers. Strikingly, CCAT1-L accumulates in cis in the nucleus while CCAT1-S is mainly cytoplasmic [13], indicating that the prevalence of altered 3' end formation of Pol II transcripts in mammals [54] may promote different IncRNP assemblies for IncRNA localization.

The coupling of IncRNA transcription and the dynamic assembly of IncRNPs during processing may also be important for both IncRNA subcellular localization and function. For example, the HAUNT IncRNA locus appears to regulate nearby HOX gene expression through the HAUNT genomic DNA, the nuclear-retained HAUNT IncRNA [55], and the transcription of this locus [56]. It will be of interest to examine how the functional outputs of many other IncRNA loci are linked to their own transcription, and whether an alteration of Pol II transcription elongation rate would affect the assembly and subcellular fates of IncRNPs and their potential roles in gene regulation.

#### Association with Specific Proteins Traps IncRNAs in the Nucleus

Mature IncRNAs may interact with nuclear matrix proteins that would subsequently facilitate nuclear localization of IncRNAs (Figure 3). The matrix protein hnRNP U is required for the accumulation of IncRNA X-inactive specific transcript on the inactive X chromosome [57] and for the in cis accumulation of FIRRE [19], which is essential for their functions. Specifically, embryonic stem cells lacking hnRNP U expression fail to form the inactive X chromosome [57] and hnRNP U is essential for FIRRE-mediated trans-chromosomal associations at its transcriptional site [19]. In addition, knockdown of hnRNP U even led to a translocation of FIRRE into the cytoplasm [19], indicating that hnRNP U is an important factor for both focal localization and nuclear retention of FIRRE. It remains to be investigated whether hnRNP U, or other unknown nuclear matrix proteins, can serve as common factors that are required for in cis accumulation or nuclear retention of IncRNAs. Such proteins would have the potential to be essential for both IncRNA subcellular localization and function. Indeed, IncRNAs might be retained in the nucleus through the formation of a mature and functional IncRNP complex including chromatin-modifying complexes or transcription factors. For instance, the interaction of nucleus-localized chromatin-modifying complexes with HOTAIR [21,22] and transcription factors with BCAR4 [25] may in turn keep these IncRNAs from nuclear export.

The higher-order organization of chromosomes may also promote nuclear retention of lncRNAs by specifying their target sites distantly. For instance, the IncRNA HOTTIP is expressed from the 5' end of the HoxA locus; it is brought to the HoxA distal genes by chromosomal looping to activate its target genes at a distance through binding the WDR5/MLL complex [58]. One speculation as to how IncRNAs are relocated to distinct sites in the nucleus is that the dynamic remodeling of IncRNPs during the lifetime of an IncRNA together with the 3D chromosomal configuration may facilitate this nuclear transportation. Future studies using genome-editing technologies to knock-in RNA aptamers at the endogenous IncRNA locus of interest followed by live cell imaging may provide substantially novel insights into IncRNA or IncRNP transport within the nucleus.



# Nuclear Retention Signals Embedded within IncRNAs Constrain IncRNAs in

The IncRNA itself may possess essential primary sequences, or more likely secondary or tertiary structures, that are required for nuclear localization (Figure 3). A nuclear retention element containing a U-rich internal loop that interacts with the 3' poly(A) tail to stabilize nuclear-accumulated RNAs has been identified near the 3' end of the nuclear-restricted polyadenylated PAN IncRNA from Kaposi's sarcoma-associated herpesvirus [59,60]. A similar structure is in the nuclear-retained MALAT1 and the long isoform of NEAT1 [60,61], but it is undetermined whether such an element acts for nuclear retention of these IncRNAs. Furthermore, a pentamer RNA motif residing in the IncRNA BORG is important for its proper nuclear localization [62]. Similarly, FIRRE has a unique highly structured 156-bp repeat named the repeating RNA domain (RRD), and FIRRE lacking the RRD fails to accumulate on the chromosome [19]. Intron-containing circular RNAs [63,64] and intron-derived sno-lncRNAs [20] are retained in the nucleus, indicating that intronic sequences may serve as ribonucleic nuclear retention elements (RNREs) that lead to IncRNA nuclear localization. Finally, some divergently transcribed IncRNAs can form DNA-RNA triplexes to anchor the IncRNAs to the vicinity of their gene loci [17].

These studies together reveal that RNA sequences can act as RNREs that mediate nuclear localization. Remarkably, such RNREs are diverse in terms of their length and sequence conservation and how RNREs promote nuclear retention of IncRNAs remains elusive. Due to the low sequence conservation of many IncRNAs, the structural domains of only a few IncRNAs have been solved. One example is the IncRNA Gas 5, which inhibits the transcriptional activity of steroid receptors (SRs) through direct competition for DNA binding. A putative 33 base stem-loop region of Gas5 has been shown to 'mimic' genomic SR-binding sites, therefore forming a compact IncRNA (Gas5)-protein (SR) interaction domain [65]. A more recent study using chemical probing and phylogenetic analysis revealed that the structural organization of HOTAIR was comparable to well-folded RNAs, like Group II introns and Gas5. The secondary structure of HOTAIR is composed of four independently folding modules corresponding to its predicted protein-binding domains [66]. It is unclear whether such folding modules can be served as RNREs for HOTAIR.

Finally, multiple factors including RNREs, higher-order chromosome organization, nuclear protein factors, and the coupling of RNP assembly with IncRNA transcription may often function together to achieve the unique subcellular localization of specific IncRNAs. The application of different approaches (Box 2) may be capable of revealing the mechanism that links IncRNA localization and function.

#### IncRNAs in the Cytoplasm

Since cytoplasmic IncRNAs and mRNAs appear similar in many aspects, an obvious hypothesis is that the export of such IncRNAs is regulated by mechanisms similar to those that regulate mRNA export, although the associated proteins for IncRNAs and mRNAs could be distinct. For example, the association of exon junction complexes (EJCs) with IncRNAs containing splice junction sites could be substantially different from that of mRNAs, as EJCs are known to be deposited on newly spliced mRNAs in the nucleus and coordinate several downstream steps of mRNA biogenesis, including nuclear export and translation (see reviews [51,67]). Interestingly, if EJCs are deposited on spliced cytoplasmic IncRNAs, then it is likely that cytoplasmic IncRNPs are not remodeled like mRNPs, which are changed dramatically by the act of translation [67]. For example, cytoplasmic IncRNAs do not go through the pioneer round of translation and they may not be remodeled the same way that mRNPs are. If true, then cytoplasmic IncRNPs may have unique and identifying RNP structures that would allow some lncRNAs escape from polyribosome trapping and mRNA turnover pathways.



#### Box 2. Emerging Approaches to Study Long Noncoding RNA (IncRNA) Localization and Function.

Visualization of IncRNAs in fixed cells is readily accomplished through regular or single-molecule RNA fluorescence in situ hybridization [10]. A new method that examines endogenous RNA in living cells uses the nuclear-localized RNA-targeting CRISPR-Cas9 system [82]. Application of native elongating transcript sequencing, which aims to study the complex events underlying transcription in mammalian cells [83], may uncover transcriptional differences between IncRNA genes that have different subcellular fates. RNA antisense purification methods enable the analysis of IncRNA-associated proteins, RNAs, and DNAs with much higher resolution. These methods have provided new mechanisms of X chromosome inactivation by identifying XIST-interacting genome architectures and proteins [84-86] and have also defined new genomic targets of NEAT1 [35] and MALAT1 [35,87]. Detailed studies are still needed to dissect which proteins are necessary for the localization or function of each IncRNA. Deciphering the structural domains of each IncRNA will be crucial to solving the specificity of long noncoding ribonucleoprotein (IncRNP) interactions. A number of reagents have been used to probe RNA structures both at the individual level in a conventional manner and at the genome-wide level through combination with deep sequencing (see review [7]). However, such studies are still far from sufficient to understand the precise molecular basis of how individual structural motifs within IncRNPs act. New methodologies and algorithms may enable the solution of structures of defined RNA structural domains with their interacting proteins. For example, small-angle X-ray scattering (SAXS) is increasingly used to characterize the structure and interactions of macromolecules in solution. SAXS has revealed that the HIV Rev response element, a 232-nt-long RNA motif that the virus uses to recognize its own mRNA among more abundant host cellular RNAs, has an unusual topological structure [88]. Electron cryomicroscopy has also recently been shown to have a powerful capability to reveal the structures of large RNA-protein complexes captured in various states, which has provided insights into yeast spliceosome assembly and catalysis [89,90]. The application of similar approaches to study IncRNP complexes will dissect the essential structures of IncRNAs and their interacting protein complexes in detail.

#### Outstanding Questions

How is the specific localization of long noncoding RNAs (IncRNAs) achieved and regulated?

Does the dynamic assembly of long noncoding ribonucleoproteins determine their subcellular localization?

How do the RNA structural motifs and effective protein partners work together to define the cellular fates of IncRNAs?

What is the life cycle of IncRNAs in the cvtoplasm?

What approaches could be applied to resolve IncRNA structural domains and IncRNA-protein complexes in detail?

#### **Concluding Remarks**

The IncRNAs are a large and diverse class of transcripts that affect gene regulation through a variety of mechanisms. Depending on their genomic origin, subcellular localization, or functional pathways, IncRNAs can be classified into different groups. Like proteins, IncRNAs must localize to specific subcellular compartments to execute their functions. However, how the specific localization of IncRNAs is achieved and regulated and what rules IncRNAs follow to make them so remarkably different from mRNAs remain largely unknown. In addition to the unexamined dynamic assembly of IncRNPs during transcription and processing, other factors including higher-order nuclear organization, RNA structural motifs, and protein partners are also proposed to play important roles in defining IncRNA subcellular localization. It will be of great interest to identify common RNA structural domains and protein factors that regulate the nuclear retention or accumulation of IncRNAs. In addition, the nucleocytoplasmic export of cytoplasmic IncRNAs and their life cycle in the cytoplasm also require a thorough investigation. Nevertheless, understanding these features of IncRNAs will greatly expand our current knowledge of IncRNA biology and shed new lights into the study of their cellular roles in depth (see Outstanding Questions).

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