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Structure and function of long noncoding RNAs in epigenetic regulation

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Genomes of complex organisms encode an abundance and diversity of long noncoding RNAs (lncRNAs) that are expressed throughout the cell and fulfill a wide variety of regulatory roles at almost every stage of gene expression. These roles, which encompass sensory, guiding, scaffolding and allosteric capacities, derive from folded modular domains in lncRNAs. In this diverse functional repertoire, we focus on the well-characterized ability for lncRNAs to function as epigenetic modulators. Many lncRNAs bind to chromatin-modifying proteins and recruit their catalytic activity to specific sites in the genome, thereby modulating chromatin states and impacting gene expression. Considering this regulatory potential in combination with the abundance of lncRNAs suggests that lncRNAs may be part of a broad epigenetic regulatory network.

Global transcriptional analyses have revealed that the vast majority of the human genome is dynamically and differentially transcribed to produce a range and complexity of lncRNAs¹. These observations are supplemented by an increasing number of targeted functional studies showing that lncRNAs fulfill regulatory roles at almost every stage of gene expression, from targeting epigenetic modifications in the nucleus to modulating mRNA stability and translation in the cytoplasm (Box 1). As a result of these studies, lncRNAs are being increasingly accepted as a major new gene class.

The abundance of lncRNAs, in conjunction with these emerging functional insights, has fuelled considerable excitement and enthusiasm for research into lncRNA biology. As the assumption of nonfunctionality has been discarded, researchers are starting to appreciate the potential importance of lncRNAs in the ontogeny of complex organisms². Research into lncRNAs has progressed so rapidly that it is becoming increasingly difficult to comprehensively catalog the functionally validated cases³. In this wide and diverse emerging functional landscape, we consider features of lncRNA structure, expression, evolution and function with respect to one of the currently best characterized role of lncRNAs—the regulation of epigenetic dynamics.

Defining IncRNAs

The majority of characterized lncRNAs are generated by the same transcriptional machinery as are other mRNAs, as evidenced by RNA polymerase II occupancy and histone modifications associated with transcription initiation and elongation⁴. These lncRNAs have a 5' terminal methylguanosine cap and are often spliced and polyadenylated. Alternate pathways also contribute to the generation of known lncRNAs, which include a poorly characterized contingent of

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non-polyadenylated lncRNAs likely expressed from RNA polymerase III promoters^{5,6} and lncRNAs that are excised during splicing and small nucleolar RNA production⁷.

Because lncRNAs have a biogenesis pathways in common with mRNA and other noncoding RNA classes, no defining biochemical features can be exclusively ascribed to lncRNAs. Rather a lack of defining features, such as the lack of an extended open reading frame (ORF), provides theoretical evidence that many transcripts function intrinsically as an RNA⁸. Conservation of an extended ORF, particularly when nucleotides in the ORF's codons exhibit different rates of selective constraint, can be used to distinguish coding transcripts⁹. Exceptions to these assumptions result from short or noncanonical peptides encoded in transcripts that evade screening attempts for viable ORFs¹⁰. Empirical support against the categorization of an lncRNA can be provided by matching ribosome footprints or peptide fragments from mass spectrometry that indicate translation^{11,12}. However, although the ability to encode a protein does not necessarily preclude a transcript from having a function as an RNA—and indeed there is a growing catalog of bifunctional mRNA that are also lncRNAs¹³—the demonstration of function as an RNA may be ultimately required for annotation as a lncRNA.

The dynamic evolutionary interface between coding and noncoding components of the transcriptome also obscures a clear annotation of lncRNAs. Coding transcripts can lose their ability to encode a protein, and noncoding transcripts can acquire a coding function^{14–16}. The complexity of this interface is seen at the X-inactivation center, where the *Xist* gene resulted from 'pseudogenization' of an ancestral protein-coding gene conserved in vertebrates, combined with the integration of flanking repetitive mobile elements¹⁷. These events remodeled the structure and sequence of the X-inactivation center in the eutherian lineage to generate noncoding transcripts that include not only *Xist*, but also *Tsix*, Jpx (*Enox*), Xite (Rr18) and Ftx (Thcytx) lncRNAs, each of which have acquired individual roles in X-chromosome inactivation¹⁸.

Alternative splicing can also incorporate exons from multiple coding and noncoding genes, thereby merging gene structures and generating ambiguous transcripts that eschew simple classification ^{1,19,20}. Such examples of transcriptional complexity have contributed to a revised concept of the transcript being the basic unit of genome

BOX 1 Cytoplasmic IncRNAs

A substantial proportion of IncRNAs reside within, or are dynamically shuttled, to the cytoplasm where they regulate protein localization, mRNA translation and stability. For example, the NFAT transcription factor is trafficked from the cytoplasm to the nucleus to activate target genes in response to calcium-dependent signals. A IncRNA, *NRON*, complexes with importin- β proteins and regulates the trafficking of NFAT¹¹³. Notably, *NRON* inhibits the trafficking of NFAT to the nucleus specifically, with other proteins also trafficked by importin- β proteins, such as NF- κ B, being unaffected.

By virtue of their ability to base pair with mRNAs, cytoplasmic IncRNAs also can regulate translation. The *UCHL1* mRNA is complemented by an antisense IncRNA, which, in response to stress or the mTOR pathway, is shuttled to the cytoplasm where, via an antisense complementary to the *UCHL1* AUG initiation codon and combined inverted SINEB2 domains, increases UCHL1 protein synthesis⁵⁴.

Additional repeat elements common to IncRNAs and mRNA create a broad interface for complementary interactions. Alu elements in cytoplasmic IncRNA can form imperfect complementary RNA duplexes with Alu elements in the 3′ UTRs of target mRNAs⁸³. Staufen1 subsequently recognizes and binds the resultant dsRNA elements and initiates mRNA decay.

expression, with the concept of a gene encompassing a hierarchy of transcripts that underlie a given phenotype²¹. Similarly, many functional precedents associated with lncRNAs can be feasibly ascribed to other transcriptional elements. For example, untranslated regions (UTRs) of mammalian mRNAs often range over kilobases, sometimes dwarfing the size of the upstream open reading frame²². Given their common chemistry, such noncoding regions can feasibly mediate functions that are similar to those of other lncRNAs²³. Although this review is constrained to the narrow definition of lncRNAs as long transcripts that exclude small regulatory RNAs, such as microRNAs, Piwi-interacting RNAs and small nucleolar RNAs, this arbitrary definition does not capture the nuanced and complex nature of the transcriptome, and efforts should be supported to evolve the concept of noncoding RNAs to a more inclusive definition of functional RNAs.

Abundance of IncRNAs encoded in the genome

Initial large-scale sequencing of cDNA libraries have revealed an unexpected abundance of lncRNAs¹. This was supported by chromosome-wide and genome-wide tiling arrays and RNA sequencing that show the human genome to be prevalently transcribed into lncRNAs^{5,24,25}. It is difficult to gauge an exact number of human lncRNAs, with current lncRNA catalogs ranging between 5,000 and 15,000 transcripts²6,27. However, there is little overlap between these different lncRNA catalogs, and this may merely represent a lower bound, with many lncRNAs yet to be annotated. Whereas the number of known human protein-coding genes has remained stable over recent years, the number of known lncRNAs continues to accumulate, and lncRNAs may eventually rival protein-coding genes in number and diversity.

LncRNAs are expressed in lower amounts generally compared to their protein-coding counterparts, making it difficult to robustly detect and assemble complex transcript structures^{26,27}. Indeed, targeted capture and RNA sequencing of intergenic regions affords the detection and assembly of many additional lncRNAs that are expressed in amounts too low to be otherwise detected by conventional high-throughput RNA sequencing (RNA-seq)²⁸. Extrapolated across the genome, this abundance of lncRNAs represents a sizable expansion of the transcriptome. Though consistent with the many regulatory functions assigned to lncRNAs, the low expression may restrict these lncRNAs to subtle or redundant roles, or reflect incomplete repression in nonspecific cells. By comparison to protein-coding genes, lncRNA expression is considerably more cell type–specific; thus, RNA sequencing for more developmental stages and tissue types will be required to achieve comprehensive annotations^{26,27}.

Organization of IncRNA loci in the genome

The genome has a modular architecture with any single sequence incorporated into a range of sense and antisense, interwoven coding

and noncoding transcripts^{1,29}. The combinatorial application of alternative splicing, transcription initiation and termination exploits this modular architecture to drive diversification of transcription³⁰. This transcriptional complexity is notably apparent in the myriad isoforms of lncRNA genes. LncRNAs are also often organized in close association with protein-coding genes (**Fig. 1a**). More than half of mammalian coding genes have complementary noncoding antisense transcription³¹, which is also accompanied by overlapping, intronic and bidirectional noncoding transcription³⁰. The iterative combination of these organizational modes generates complex transcriptional loci that include both coding and noncoding transcripts¹ (**Fig. 1b**).

Although complex loci containing lncRNAs and protein-coding gene may evolve to have a common local regulatory architecture, this clustered organization may also reflect the cis-acting functions of many lncRNAs in mediating changes to the local chromatin and the expression of neighboring genes. The X-inactivation center illustrates how the architecture of a complex locus regulates expression of the central Xist transcript (Fig. 1c). The lncRNA Tsix is transcribed antisense to, and initiates silencing of, *Xist* expression from the active X chromosome³². Tsix transcription through the 5' end of Xist establishes a repressive chromatin domain by interaction with Polycomb repressive complex 2 (PRC2) and enhancing hypermethylation via DNA methyltransferase 3A (DNMT3A), thereby mediating long-term *Xist* silencing^{33,34}. However, neither DNA methylation nor PRC2 is required for Xist repression³⁵, indicating that Xist regulation by Tsix is complex, with additional and alternative pathways. Indeed, the lncRNAs Tsix and Xist can form long RNA duplexes that are subsequently processed by the RNA interferences pathway into small regulatory RNAs that can contribute to downstream epigenetic changes³⁶. Notably, *Tsix* has an additional antisense partner, Xite (Rr18), that also functions as a cis-acting activator of Tsix, appending additional layers to this regulatory circuit³⁷.

RNA structure and chemistry dictate IncRNA function

Given that the defining characteristic of lncRNAs is an intrinsic ability to function as an RNA molecule, it is important to understand the features of RNA chemistry from which this functionality derives. In addition to the four core nucleotides, the RNA sequence can include more than 100 chemically distinct modified nucleotides³⁸. To date, most modifications have been detected in tRNAs, rRNAs and small nucleolar RNAs, where they often modulate and stabilize RNA structures³⁹. However, the conversion of modified nucleotides for detection with high-throughput sequencing is revealing widespread nucleotide modifications throughout the transcriptome⁴⁰, where the modifications may similarly modulate lncRNA function^{41,42}. Many of these post-transcriptional modifications are reversible and, given the range of modifications and targets, may comprise an additional

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Figure 1 The human genome encodes an abundance and diversity of IncRNAs. (a) LncRNAs can be found harbored in intergenic regions or often clustered with protein-coding genes in complex transcriptional loci (top). Schematic of transcriptional networks (bottom) shows examples of IncRNAs (red) organized bidirectionally, or antisense to proteincoding genes or in introns of protein-coding genes (blue). Alternative splicing generates many IncRNAs isoforms and can merge gene structures by incorporating both coding and noncoding exons into a single transcript. (b) The X-inactivation center illustrates a complex IncRNA locus with numerous overlapping IncRNAs¹⁷, including *Tsix*, *Ftx* and *Jpx*, that together regulate the expression of Xist, which inactivates the female X chromosome. (c) An example of the complex organization of IncRNAs associated with developmental genes. The Sox2 gene involved in pluripotency and development is surrounded by many overlapping IncRNAs¹¹².

layer of post-transcriptional regulation analogous to the epigenetic landscape⁴³.

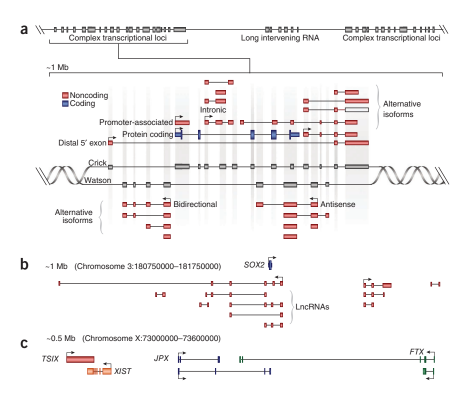
A major feature of lncRNAs is a propensity to fold into thermodynamically stable secondary and higher-order structures. RNA has the

capacity to form hydrogen bonds on the Watson-Crick face but also the Hoogsteen and ribose face. These collective interactions contribute to RNA secondary structures that include double helices, hairpin loops, bulges and pseudoknots, and that are connected in higher-order tertiary interactions primarily mediated by non–Watson-Crick base-pairing⁴⁴. This results in an RNA architecture dominated by coaxial stacks of helices that are organized in parallel or orthogonal to one another. This architecture is also modular, with recurrent structural motifs, including the sarcin-ricin loop, the K-turn and the C-loop, that exhibit only minor dependencies on neighboring sequences⁴⁵. Furthermore, given that Watson-Crick base pairing and base stacking provide the greatest contribution to the energetic stability of RNA structures, these modular secondary structures generally fold initially and independently, before subsequent tertiary interactions occur, resulting in the hierarchal assembly of RNA structure⁴⁶.

In many cases, the secondary structure of lncRNAs dictates their function. For example, conservation of the secondary structure maintains the tumor suppressor function of lncRNA MEG3, rather than its primary sequence⁴⁷. Although most RNA structural motifs originally had been described from rRNA and tRNA genes⁴⁸, increasing attention has since focused on empirical determination of lncRNA structure. Recently, the human lncRNA SRA1, a coactivator for several hormone receptors, was subjected to detailed chemical and enzymatic probing to determine four broad domains that encompass a suite of secondary structures⁴⁹. Combining enzymatic and chemical probing with sequencing enables high-throughput characterization of RNA secondary structures required to delineate structure-function relationships^{50,51}. These initial studies have confirmed a complex structural landscape in lncRNAs that can be distinguished from mRNAs on the basis of their high folding energy⁵¹.

Range of functional domains in IncRNAs

To dissect the functional structures and sequences in lncRNAs, we can borrow insight and terminology developed in the field of synthetic biology, where RNA is commonly used as a regulatory device



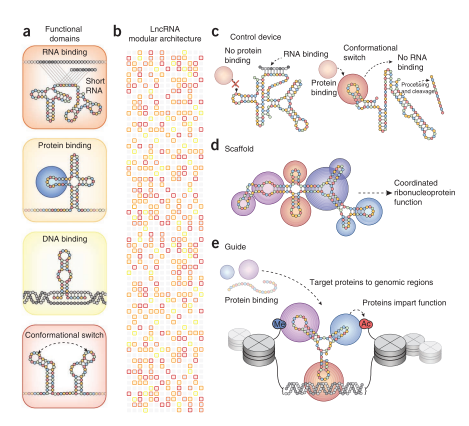
in genetic circuits⁵². RNA is a preferred substrate for such devices because it can rapidly shift between multiple stable structural conformations and undergo allosteric transitions, thereby acting as a responsive switch. Owing to the omission of translational processes, synthetic or noncoding RNAs are processed faster than other molecules dependent on transcription- or translation-dependent processes, a compact genetic footprint and a reduced energetic and resource load on the host cell⁵³. These advantages of synthetic RNA regulatory devices similarly apply to lncRNAs.

RNA-binding domains. By virtue of their ability to base pair with other RNAs, lncRNAs can act as highly specific sensors of mRNA, microRNA and other lncRNA expression (Fig. 2), and RNA aptamers are often designed to induce a programmable response in riboswitches⁵². Antisense lncRNAs can regulate the stability and translation of complementary mRNAs. For example, translation of the *UCHL1* mRNA is regulated under stress by action of an lncRNA with antisense sequence that is complementary to, and encompasses, the *UCHL1* start codon⁵⁴.

Although antisense lncRNAs are prevalent in the genome, it is notable that small and often imperfect regions of nucleotide complementarity are sufficient for specific interactions, as demonstrated by the potent ability of microRNAs to target a select suite of mRNAs via short, imperfect seed sequences⁵⁵. This imperfect complementarity has the advantage of allowing multiple RNA agonists, each with a range of dissociation constants, to compete for binding, thereby permitting lncRNAs to sense disparate RNA expression signals in the cell. For example, imperfect base pairing between Alu elements in lncRNAs and the 3′ UTR of translationally active mRNAs results in a dsRNA structure recognized and bound by Staufen1 and subsequently targeted for degradation.

The cleavage of lncRNAs can also generate small RNAs that serve as an output signal⁵⁶. A small tRNA-like sequence is cleaved from the 3' end of the lncRNA MALAT1 and trafficked from the nucleus to the cytoplasm⁵⁷. Similarly, the formation of extended RNA duplexes or

Figure 2 Domain architecture of IncRNAs. (a) LncRNAs contain structural domains that can sense or bind other RNAs via complementary base pair interactions, proteins and possibly DNA that can induce allosteric conformational changes to other structures in the IncRNA. (b) Alternative splicing can combine these structural domains into the modular architecture of IncRNAs. Each row represents an individual IncRNA; colors correspond to those in a. (c) Coupling sensory and actuator domains permits IncRNAs to act as a control device. In the example on the left, the binding of RNA (gray) induces a conformational change that prevents protein binding. Alternatively, as illustrated on the right, the protein can bind in the absence of the RNA, inducing the formation of a stem-loop secondary structure that can be processed and cleaved to generate an RNA output. (d) LncRNAs such as HOTAIR can act as molecular scaffold by binding multiple proteins to form complex ribonucleoprotein structures⁷⁸. (e) LncRNAs, such as Xist, can target the catalytic function of proteins to specific sites in the genome⁷⁹. LncRNAs can recruit chromatin-modifying proteins (purple and blue) to target sites by association with a DNA-binding protein such as YY1 (red) 71 . The chromatin modifiers then modify local histones to influence the expression of adjacent genes.



stem loops provides a ready substrate for Dicer enzyme to generate multiple small regulatory RNAs that have cascading ability to mediate downstream epigenetic changes⁵⁸. Ribozymes comprise RNA secondary structures capable of the phosphodiester bond cleavage within themselves or in other RNAs⁵⁹. Comparison between long and short RNA populations in human cells suggests widespread evidence of post-transcriptional cleavage, with lncRNAs being a preferred substrate for the generation of small RNAs²¹. The use of RNA as both output and input signals promotes RNA as a standard medium for transferring information within and between regulatory pathways, thereby assembling complex, multilayered and modular regulatory networks in the cell.

Protein-binding domains. Proteins are a major partner of lncRNAs (Fig. 2), with complexed ribonucleoprotein (RNP) particles acting as chaperones, transport aids or effectors⁶⁰. Although RNA-binding proteins are one of the most abundant human protein classes, they are assembled from relatively few RNA-binding modules, and these domains are deployed in modular combinations with intervening disordered linker regions to accommodate the large diversity of RNA structures^{61,62}. Additionally, many noncanonical domains supplement these RNP catalogs, with kinases, DNA-binding proteins and metabolic enzymes also found bound to RNA⁶³, and analysis of RNP fractions estimated that at least ~15% of expressed proteins are associated with polyadenylated RNA⁶⁴.

Approaches such as photoactivatable ribonucleoside-enhanced crosslinking and immunoprecipitation (PAR-CLIP) provide the complementary footprint to protein interactions across the transcriptome⁶⁵, and details of such protein-RNA interfaces can be resolved by X-ray crystallography or NMR spectroscopy. Proteins tend to interact with RNA where it forms complex secondary structures, positioning protein structures into the groove of an RNA stem-loop helix or providing a binding pocket in β -sheets for unpaired RNA nucleotides⁶⁶. Almost all such interactions characterized to date involve conformational changes to the protein, the RNA or both. The structural diversity of RNA in combination with the commensurate abundance of RNA-binding proteins provide a broad interface for communication between the proteome and transcriptome.

DNA-binding domains? There is currently little evidence for direct interaction between lncRNAs and DNA. RNA:DNA hybrids or triplex structures can allow single strands of RNA to interact with DNA duplexes by base-pair interactions. These direct RNA-DNA interactions could efficiently and selectively target RNA signals to genomic loci (Fig. 2). However, such interactions may also expose the genome to deamination and damage^{67,68}. A promoter-associated lncRNA, pRNA, can occlude binding of the transcription termination factor 1 (TTF1), while simultaneously recruiting DNMT3b to repress rRNA gene expression⁶⁹. Notably, this lncRNA can form a triplex with the TTF1-binding site in vitro, supporting a direct interaction with the genomic locus. Rather than being involved in direct complementary interactions, RNA folds may create a DNA-binding pocket in a manner analogous to the DNA-binding domains of a protein transcription factor. Similar to protein transcription factors, an enriched DNA sequence motif has been identified in the binding sites of an lncRNA, HOTAIR⁷⁰. Alternatively, rather than directly interacting with DNA, Xist harnesses the sequence-specific YY1 transcription factor to tether Xist to sites in the X chromosome 71 .

Modular architecture of IncRNAs

LncRNAs can act as regulatory devices by allosterically coupling binding domains with the switching of structural conformations and thereby activating or suppressing linked functional domains⁵². Incorporating multiple sensors into the architecture of a single lncRNA permits the integration and processing of multiple inputs through logic gates to produce a single output⁵³. An instructive example of such a switch embedded in an RNA structure involves a protein-dependent riboswitch in the VEGF 3' UTR that is bound by either GAIT or heterogeneous nuclear ribonucleoprotein L (hnRNP L) protein complexes 72 . The binding of one protein, in response to inflammatory and hypoxic cues, induces a conformational change that precludes binding of the other protein, thereby processing different protein inputs to regulate mRNA translation and gene expression.

When each of these modular domains specifies alternate protein interactions, an lncRNA can scaffold many disparate proteins into a single cogent ribonucleoprotein⁷³. The lncRNA CDKN2B-AS1 (ANRIL) binds the Suz12 subunit of the PRC2 complex and the chromodomain of CBX7 in the PRC1 complex to trimethylate H3K27, which then mediates repression of the CDKN2A/CDKN2B (IFNK4a/ IFNK4b) tumor suppressor locus⁷⁴. With at least 12 chromatinmodifying proteins having been associated with lncRNAs to date⁷⁵, the composition of possible chromatin-modifying proteins in a single ribonucleoprotein can be varied by shuffling the modular components within an lncRNA, in concordance with the combinatorial histone code⁷⁶. For example, in mice the lncRNA Kcnq1ot1 can recruit both PRC2 and G9a, which impart H3K4 trimethylation and H3K9 methylation, respectively⁷⁷. Similarly, HOTAIR binds and targets the catalytic methyltransferase subunit EZH2 of PRC2 via a structural domain at the 5' end of HOTAIR. The 3' end binds and recruits lysine-specific demethylase 1 (LSD1), which catalyzes demethylation of H3K4, a modification similarly associated with gene repression⁷⁸. HOTAIR can therefore act as a scaffold that, via its modular architecture, can combine PRC2 and LSD1 into a single RNP complex.

Many functional domains in lncRNAs derive from repeat elements. Xist exhibits such a modular functional architecture, containing an A-region comprised of up to nine repetitive elements that fold into two stem-loop structures that recruit PRC1 for H3K27 trimethylation, and a C-repeat region that binds to both hnRNP U and YY1 and is responsible for localizing and tethering Xist to the X chromosome^{71,79-81}. Each of these functional domains, the A- and C-elements, derives from the insertion and remodeling of repetitive elements. This repetitive architecture is also present in the functionally analogous lncRNA Rsx, which mediates X-chromosome inactivation in marsupials. Although Rsx likely evolved independently and has no homology with Xist, Rsx is similarly enriched for tandem repeats in the 5' region and conserved stem-loop motifs⁸². Similar repetitive sequences identified across many lncRNAs may also reflect underlying functional domains in lncRNAs, analogous to protein domains²⁶. Transposon-derived repeats, such as short interspersed elements (SINEs) and Alu elements, can also be integrated within lncRNAs, where they act as recognition domains that target complementary mRNA for translation or degradation^{83,84}. In these cases, the transmission of transposons can result in the insertion of a ready functional domain into target lncRNAs.

Cis and trans mechanisms of epigenetic regulation

The regulatory potential of lncRNAs has been best characterized in the context of their role as epigenetic modulators. During development, DNA and histone modifications comprise a detailed and dynamic template that is overlaid across the genome to guide the regulation and expression of the underlying DNA sequences. It has become apparent that lncRNA can guide many of the proteins that catalyze these modifications^{75,76,85,86}.

The lncRNA HOTAIR guides chromatin proteins and their catalytic action in *trans* to multiple sites spread across the genome. *HOTAIR* is expressed from the end of the *HOXC* (*HOXC*@) cluster in cells with distal and posterior identities⁸⁷. HOTAIR binds and targets PRC2

to the *HOXD* (*HOXD*@) cluster as well as hundreds of additional sites around the genome to impart repressive histone modifications⁷⁰. These focal interactions of HOTAIR with target genome sites are likely pioneering events that subsequently nucleate broad regions of Polycomb occupancy and H3K27 trimethylation. By the expression of *HOTAIR*, distal developmental states can initiate an epigenetic regulatory cascade that maintains the cells' positional identity and continually refines a progressive developmental trajectory.

Many lncRNAs also mediate local functions in *cis*, recruiting chromatin-modifying proteins to modify their surrounding epigenetic neighborhood. For example, loci with parent-of-origin-specific expression patterns, termed imprinted regions, often encompass many lncRNAs^{86,88}. These lncRNAs can silence expression of neighboring genes on the parental allele from which they are expressed. Air (Airn), an 108-kb lncRNA, is transcribed from the paternal allele to which it recruits G9a to methylate H3K9 residues over an adjacent 300-kb region of the genome, thereby silencing the expression of neighboring paternal *Igf2r*, *Slc22a2* and *Slc22a3* genes⁸⁹. The ability of lncRNAs to act in *cis* to remodel the local epigenetic landscape and thereby regulate neighboring gene expression may, in part, explain the concentration of lncRNAs around many critical developmental genes whose expression is subject to strict control⁹⁰ (**Fig. 1c**).

Local epigenetic changes initiated by lncRNAs can seed broader epigenetic effects, with the silencing of an entire female X chromosome by Xist providing a dramatic example. After transcription, Xist accumulates at many sites across the X chromosome that subsequently seed the progression of facultative heterochromatin formation ^{91,92}. LncRNAs similarly seed the spread of heterochromatin at pericentric satellites ⁹³. Pericentric satellite repeats, localized around the centromere, are required for genome integrity and replication. The peripheral regions of these satellite repeats undergo bidirectional transcription during early development, and the resulting forward-strand lncRNAs can recruit heterochromatin protein 1 (HP1) proteins to the satellite repeats to impart a distinct modified chromatin profile ⁹⁴. The continued local accumulation of HP1 results in the spreading of facultative heterochromatin to demarcate broad repressive chromosomal domains.

Role for IncRNAs in nuclear organization

Considering complex transcriptional loci and their surrounding epigenetic landscape in the three-dimensional context of the nucleus can obscure the distinctions between *trans*- and *cis*-acting lncRNA functions. The HOTTIP lncRNA binds WD repeat-containing protein 5 (WDR5), a cofactor to the histone methyltransferase MLL1 that mediates H3K4 trimethylation associated with gene activation⁹⁵. Although HOTTIP remains bound at the site of transcription, chromatin looping positions the HOTTIP-WDR5-MLL1 complex into close spatial proximity to multiple distal sites to coordinate active chromatin domains and gene expression throughout the *HOXC* loci. Chromosome looping thereby extends the range to which HOTTIP can impact epigenetic changes, similar to the role of looping in facilitating enhancer action over large genomic distances.

Immunofluorescence microscopy has revealed a complex internal structure to the nucleus, with modified chromatin often localized in cogent subnuclear structures. LncRNAs can scaffold and nucleate the assembly of such structures, recruiting the accumulation of proteins required for their assembly. For example, the lncRNA MALAT1 binds to and sequesters serine/arginine-rich splicing factors to nuclear speckles⁹⁶ and the lncRNA NEAT1 is a requisite structural component of interchromatin paraspeckles⁹⁷. Additional subnuclear structures, including histone locus bodies and stress



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BOX 2 The 'evolvability' of RNA?

The close relationship between nucleotide sequence and RNA and protein secondary structure has proved an ideal model system for the analysis of genotype-phenotype relationships in the context of adaptive evolution¹¹⁴. The structure (or phenotype) of RNA or proteins can be generated by a number of alternative sequences, as different RNA or protein sequences may fold into similar structures. As a group, these different sequences are called a genotype network. Generally, many sequences fold into similar RNA secondary structures, and therefore RNA structures have very large and diverse genotype networks¹¹¹. Indeed, entirely dissimilar sequences can often fold into similar RNA structures, and RNA structures can therefore traverse the nucleotide space completely. For example, RNA molecules that adopt the characteristic tRNA cloverleaf secondary structure can differ in up to >90% of their nucleotides¹¹⁵, and it is typical for naturally occurring RNAs to have astronomically large genotype networks¹¹⁶. Protein structures have much smaller genotype networks that are closely restrained in their ability to traverse the nucleotide space¹¹¹, and have a much lower affinity to explore other viable secondary structures. Random protein libraries rarely fold into soluble and compact protein structures¹¹⁸. By contrast, random RNAs collapse with high probability into compact and ordered structures¹¹⁸.

The large genotype network size of RNA structures means they are robust, capable of preserving secondary structure while accumulating mutations ¹¹⁹. However, these accumulated mutations also serve to diversify the genotype network, spreading the network into more neighborhoods and making a broader spectrum of a novel phenotype in immediate neighborhood directly accessible by only a small number of additional mutations¹¹⁹. This ultimately increases the likelihood of encountering a beneficial phenotype and therefore achieving evolutionary adaptation and innovation. This advantage is demonstrated in ribozymes that harbor greater levels of latent cryptic variation being better suited to evolutionary adaption in response to changing environment conditions¹²⁰. Collectively, this suggests that RNAs, in comparison to proteins, have a higher affinity for adaptive evolution as a result of their greater genotype networks.

The complexity of the transcriptome, driven in large part by a massive expansion of lncRNA transcripts, is a hallmark of the eukaryotic genome. It is tempting to speculate that this expansion of lncRNAs has been driven in part by selection for their rapid evolvibility, which reciprocally has been a primary factor in driving the radiation and evolution of eukaryotic lineages¹²¹.

bodies also require RNA for nucleation and assembly, suggesting a broad role for lncRNAs in nuclear organization⁹⁸.

Nuclear organization also forms an overarching three-dimensional context under which lncRNAs mediate their regulatory roles. For example, repressive regions of H3K27 trimethylation and PRC2 occupancy form a network of close physical interactions that appear as discrete Polycomb bodies in the nucleus 99,100. Given the close and promiscuous association of PRC2 with lncRNAs85, the cis-acting action of many lncRNAs may be similarly involved in mediating the repressive functions of Polycomb bodies. Indeed, two lncRNAs, TUG1 and MALAT1, traffic gene loci between Polycomb bodies to the activating context of interchromatin granules¹⁰¹. TUG1 lncRNA specifically associates with methylated PRC2 at E2F1 target gene promoters within Polycomb bodies. However, in response to growth signals, the promoter-localized PRC2 is demethylated and associates with MALAT1, resulting in the relocation of these gene loci to interchromatin granules. This interaction with MALAT1 also regulates E2F1 sumoylation, which permits recruitment of CDCA7L, a histone H2B monoubiquitinase, to switch the preference of the PRC2 chromodomain from repressive to activation-associated histone modifications¹⁰¹.

LncRNAs in a broader epigenetic regulatory network

Many of the examples discussed above ascribe a function of lncRNAs in guiding, whether in cis or trans, the catalytic function of chromatin-modifying proteins to specific genomic sites. Thousands of lncRNAs are found in association with chromatin modifiers, associations that are as highly dynamic as the tissue- and developmentspecific expression of lncRNAs^{75,85}. Considering the abundance of lncRNAs along with these functional precedents raises the potential for lncRNAs and chromatin-modifying enzymes to collectively comprise a regulatory network with the requisite complexity to delineate a dynamic epigenetic landscape. A recurrent property of biological regulatory networks is a scale-free topology, in which the majority of nodes have few links whereas a minority of nodes, termed hubs, are furnished with many links that bind the network together 102. Chromatin-modifying enzymes as ubiquitous and highly connected proteins are archetypal hubs, whereas many lowly expressed, and tissue-specific lncRNAs likely populate sparsely connected, lowerlevel nodes.

A major advantage of scale-free networks is that they are robust. The network is tolerant to the inactivation of even a large number of sparsely connected nodes without disrupting the integrity of the network, consistent with the few phenotypic effects observed in knockout screens targeting individual lncRNAs^{103–105}. However, scale-free networks are vulnerable to the inactivation of highly connected hubs that fatally splinter off the network into many isolated nodes. Indeed, chromatin-modifying proteins are highly conserved, with few genes being lost between worms and humans¹⁰⁶, and mutations in chromatin-modifying proteins causing multiple cancers and developmental disorders¹⁰⁷.

Selective constraint would also be relieved on nonessential lncRNAs nodes. LncRNAs are among the fastest evolving elements in the genome, with a rate of lncRNA gain and loss that is much higher than that of protein-coding genes^{108–110}, and one-third of human lncRNAs are thought to have arisen solely in the primate lineage²⁶. This specificity extends to function, with even the iconic HOTAIR exhibiting divergent roles in human and mouse^{27,104}. Therefore, combining chromatin modifiers and lncRNAs in a cogent regulatory network could confer both robust and adaptive capacities¹¹¹ (Box 2).

Future directions

Since the abundance of lncRNAs was revealed in early sequencing efforts²⁴, lncRNAs have been the focus of intense research, and already a wide range of functional roles have been ascribed to individual lncRNAs. However, the sheer abundance and diversity of lncRNAs pose a challenge for their characterization.

A greater understanding of structure-to-function relationships—that is, how and which modular elements dictate a specific function—will be required to characterize such an abundance of transcripts. The development of high-throughput approaches to determine secondary structure, protein-binding motifs and other features in the primary sequence could realize a detailed and global landscape of elements in lncRNAs. The functional characterization of such elements, as opposed to individual transcripts, would provide a powerful predictive platform to extrapolate functions across related classes lncRNAs that have similar features. This could ultimately permit the functional assignation and validation of many lncRNAs on the basis of sequence and structure, analogous to



306

large-scale structural proteogenomic efforts, and be hugely informative in the hypothesis of individual lncRNA function.

LncRNAs have, in a relatively short period of time, become recognized as a legitimate and major new class of genes. LncRNAs may potentially comprise a major component of the genome's information content, complementary and comparable in abundance and complexity to the proteome. Given such huge potential, they have begun to generate considerable excitement in the molecular biology community. It is with time that we will realize whether lncRNAs will live up to such potential.

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COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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