Minireview

Employment opportunities for non-coding RNAs

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Abstract Analysis of the genomes of several eukaryotic organisms, including mouse and human, has reached the striking conclusion that the mammalian transcriptome is constituted in large part of non-protein-coding transcripts. Conversely, the number of protein-coding genes was initially at least overestimated. A growing number of studies report the involvement of non-coding transcripts in a large variety of regulatory processes. This review examines the different types of non-coding RNAs (ncRNAs) and discusses their putative mode of action with particular reference to large ncRNAs and their role in epigenetic regulation.

Keywords: Non-coding RNA; Antisense transcript; Intergenic transcription; X-chromosome inactivation

1. Introduction and historical context

In the 1960s when Jacob and Monod [1] first defined the basic principles underlying bacterial genetic regulatory systems, the existence of two distinct types of genes was already clearly established: structural genes encoding mRNAs for proteins and regulatory genes producing regulatory RNAs that interact by base pairing with the operator elements of structural genes. For much of the last twenty years, at least as far as higher eukaryotes have been concerned, interest in the transcriptional activity of the genome has been focussed almost exclusively on gene discovery and protein-coding genes. Progressively, the dogma that the critical functions of the cell depend exclusively on proteins gained ground, the complexity of an organism depending exclusively on its repertoire of protein-coding genes. RNAs were considered solely as accessory molecules, involved mainly in mediating the processes of transcription and translation. This over-simplistic view was first called into question by the discovery of untranslated intronic sequences embedded within coding genes and the associated finding of splice variants which allow the synthesis of more than one protein product from a single gene (for review see [2]). Identification of the catalytic properties of the RNA subunit of ribonuclease P further underlined the notion that the functions of RNAs extend well beyond a transient role in ensuring the expression of protein-coding genes (for review see [3]).

Recent advances in genome sequencing and the availability of extensive finished sequence for the human and mouse genomes has not significantly improved our ability to identify putative regulatory RNAs. Much of the sequence related effort in genomics, which has been concentrated on developing methodologies to identify and define classical genes within genomic sequence, is based on the identification of conserved coding exons by comparative genome analysis [4] or on computational gene prediction which relies on gene-finding algorithms [5]. Such gene-finding algorithms are designed to identify open reading frames (ORFs), polyadenylation signals, conserved promoter regions and splice sites typically associated with protein-coding genes. They generally do not allow the detection of non-coding RNA (ncRNA) genes, which often show only weak primary sequence conservation, lack ORFs and are not systematically processed. ncRNAs remained cloistered until very recently, when relevant computational and experimental approaches were at last initiated. In yeast, searches for polymerase III promoters, which often characterize small RNA genes, and analysis of expression profiles within “gaps” between predicted ORFs have identified new ncRNAs. Of particular interest is a recent comparison between the sequence annotations available for human chromosomes 21 and 22 and the expression profile of these chromosomes as established by hybridization of polyA + total RNAs extracted from 11 different human cell lines using oligonucleotide arrays, which cover the entire genomic DNA sequence of these chromosomes. These studies have led to the conclusion that there are 10-fold more transcription units than predicted coding genes in the human genome [6]. A small fraction of these additional transcription units is thought to represent pseudogenes, as sequence analysis has revealed that many of the 20000 pseudogenes in the human genome possess functional promoters. (Note, however, that transcription of some pseudogenes may exert a regulatory function [7].) A larger fraction of this transcriptional activity is probably derived from repeat sequences that are transcribed such as long terminal repeats, small interspersed elements and long interspersed elements (LINEs). Many of these repeat elements contain active promoters; often containing canonical P0/II recognition sites.
Others, such as many of the LINEs are truncated at their 5′ end. Accidental insertion of such elements, even when truncated, downstream of an endogenous promoter might however be expected to result in transcription. How many of the aberrant transcripts originating from such so-called “junk” DNA are retained and how many are recognized as illegitimate and rapidly destroyed by mechanisms such as the non-sense-mediated decay pathway, remains to be defined [8].

Originally, the term ncRNAs referred exclusively to polyadenylated eukaryotic RNAs transcribed by RNA polymerase II, carrying a 7-methylguanosine cap structure and lacking an ORF. Nowadays, this definition has been extended to designate all RNA transcripts without protein-coding capacity. Such ncRNAs can be divided into two classes: housekeeping RNAs and regulatory RNAs. Housekeeping ncRNAs (Table 1) are usually small, constitutively expressed and necessary for cell viability. They include not only ribosomal RNAs (rRNAs) and transfer RNAs (tRNAs) but also small nuclear RNAs (snRNAs) and small nuclear RNAs ( snoRNAs) implicated in diverse functions including splice regulation and rRNA modification. Housekeeping ncRNAs also include RNAs important for the transport and insertion of proteins into membranes and telomeric sequence addition (for review see [9]). A subset of the additional transcription units, that have been detected in global transcriptional profiling experiments, will probably correspond to novel ncRNAs falling into this category. The recent elaboration of software based on algorithms using “covariance models”, which allow the detection of consensus secondary structures, may provide a way to identify and associate these small ncRNAs with cellular functions [10].

Regulatory ncRNAs include the microRNA (miRNA) family, which can induce post-transcriptional gene silencing (PTGS) by repressing mRNA translation (Fig. 1B). miRNAs are involved in the control of developmentally timing and/or tissue-specific functions. The first miRNAs to be described, corresponded to the products of the lin-4 and let-7 genes in Caenorhabditis elegans. These miRNAs, 22 and 21 nucleotides in length, are processed from larger precursor RNAs and inhibit translation through antisense interactions with the 3′ untranslated region of the target mRNAs. miRNAs are widely distributed in all organisms. They are encoded in the genome as short inverted repeats which have a double-stranded RNA (dsRNA) stem loop about 70 bp long [11] and are found in introns as well as in intergenic clusters. Processing produces the small of 21–25 nucleotide effector molecules, which are usually derived from only one strand of the stem loop structure. Some miRNAs have homologues in both vertebrates and invertebrates although their small size renders the criteria of conservation between species often insufficient for the identification and isolation of new miRNAs. The number of miRNAs in human is thought to be 220–250 [12].

Besides such small ncRNAs, an even greater mystery envelops the role of larger regulatory ncRNAs. These RNAs show great diversity in their genomic organization. In some cases, they are produced from within a well-defined gene (in such cases, they are processed: spliced and/or polyadenylated), whilst in others, transcription either initiates within the intron of a host gene or may result from intergenic transcription (Fig. 1A). The observation that many regulatory elements such as locus control regions (LCRs), boundary elements, silencers and insulators are transcribed, points to the probable involvement of intergenic non-coding transcription in the function of these genomic regulators [13]. ncRNAs may also be transcribed from a single strand (sense ncRNAs) as well as in the opposite orientation when they may overlap with either protein-coding or non-coding genes (antisense RNAs transcription). Many examples of cis-acting natural antisense transcripts (cis-NATs) have recently been described (for review see [14]). Two recent studies, one using the FANTOM2 mouse cDNA set, public mRNA data and mouse genome sequence data [15], the second combining a computational based search for sense-antisense transcripts pairs in human genome public databases with experimental assessment of the results using microarrays containing strand-specific oligonucleotides probes [16], have shown that the contribution of NATs has been largely under-evaluated in both mouse and human. It is now estimated that an antisense transcript is transcribed from some 10–20% of genes. Due to database bias in favour of polyA+ RNAs, most of the antisense transcripts detected in these analyses were spliced and polyadenylated. This artefactual bias in favour of processed antisense RNAs may suggest that the overall proportion of antisense transcripts may be even higher. All possible types of arrangements of cis-NATs relative to their sense counterpart have been described: tail-to-tail, head-to-head or arrangements with the antisense transcript totally included within an intron of the sense transcript have been described [15]. Antisense ncRNAs appear to be especially abundant at imprinted loci, which are usually organized in

Table 1
Functional classification of housekeeping ncRNAs

<table>
<thead>
<tr>
<th>Type</th>
<th>Function</th>
<th>Databases or search tools</th>
</tr>
</thead>
<tbody>
<tr>
<td>rRNA</td>
<td>Translation of genetic information</td>
<td><a href="http://intra.psb.ugent.be:8080/rRNA/">http://intra.psb.ugent.be:8080/rRNA/</a></td>
</tr>
<tr>
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</tr>
<tr>
<td>snRNA</td>
<td>Pre-mRNA splicing; spliceosome components</td>
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</tr>
<tr>
<td>snoRNA</td>
<td>RNA modifications, 2′-O-methylation and</td>
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</tr>
<tr>
<td></td>
<td>pseudouridylation</td>
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</tr>
<tr>
<td>tmRNA</td>
<td>trans-translation</td>
<td><a href="http://psyche.uchtd.edu/dbs/tmDB/trimRDB.html">http://psyche.uchtd.edu/dbs/tmDB/trimRDB.html</a></td>
</tr>
<tr>
<td>Telomerase RNA</td>
<td>Telomeric DNA synthesis</td>
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</tr>
<tr>
<td>Signal recognition</td>
<td></td>
<td><a href="http://psyche.uchtd.edu/dbs/SRPDB/SRPDB.html">http://psyche.uchtd.edu/dbs/SRPDB/SRPDB.html</a></td>
</tr>
<tr>
<td>Particle</td>
<td>RNA processing</td>
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<tr>
<td>Ribonuclease RNA</td>
<td></td>
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</tr>
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Non-coding housekeeping RNAs are outside of the focus of this review. Details about this class of transcripts can be found in the indicated databases.

Abbreviations: rRNA, tRNA, snRNA, snoRNA, tmRNA, transfer-messanger RNA.
clusters and are regulated by putative imprinted centres [17]. Antisense transcripts have been found for 22 of the 58 mouse imprinted genes examined [15].

One of the best-characterized regions producing different subtypes of ncRNAs and containing non-coding transcription units is the mouse X-inactivation centre (Xic) (Fig. 3), the master control region implicated in the regulation of X-chromosome inactivation (XCI) in mammals [18]. Through reference to this system and other particularly informative examples, we will focus the rest of this review on the functions potentially exerted by large regulatory ncRNAs and the associated underlying molecular mechanisms.

2. ncRNAs and the long-range establishment of high-order chromatin structures

In mammals, dosage compensation of X-linked gene products between the sexes is achieved by the transcriptional silencing of a single X-chromosome during early female embryogenesis. Initiation of XCI requires the counting of X-chromosomes and the random choice of the X-chromosome to inactivate (for review see [19]). Once initiated, the inactivation signal is propagated along the chromosome inducing its heterochromatinization. This inactive state is stably maintained through subsequent cell divisions. The Xic, which controls XCI initiation and spreading, contains the X-inactive specific transcript (Xist) gene. Xist was first discovered in 1991 and corresponds to one of the first gene described to encode an untranslated RNA [20–22]. The gene is conserved between species at the level of its genomic organization but shows only weak sequence homology, possibly implicating a role for secondary structure [18]. Xist ncRNAs are 15–17 kb long in mice, spliced, polyadenylated and restricted to the nuclear compartment [23,24]. Prior to the onset of XCI, Xist RNA is synthesized from both X chromosomes in the female embryo. Xist up-regulation on the putative inactive X-chromosome and RNA coating of this chromosome constitute the first detectable signs of inactivation. This coating and Xist upregulation then trigger the enrichment of the targeted X by the lysine-27 methylated form of histone H3 (H3K27). The establishment of this modification is ensured by the polycomb group complex eed enxl, which possesses a histone H3K27 methyltransferase activity and which transiently accumulates on the future inactive X at the time of XCI [25,26]. H3K27 hypermethylation is accompanied by other chromatin changes, including H3K4 hypomethylation, H3K9 hypermethylation and H3–H4 hypoacetylation. These modifications appear concomitantly with
the transcriptional silencing of X-linked genes. It remains unclear whether this silencing is directly triggered by Xist accumulation or is subsequent to chromatin modifications. Replication timing of the inactive X then becomes retarded into late S-phase, whilst the CpG islands associated with the promoters of X-linked genes are methylated [27]. These numerous and successive layers of modification lead to the establishment of a silent chromatin structure on the elected X and, in turn, lock the inactive X into a stable heterochromatic state, which ensures the faultless maintenance of the inactive state throughout the cell cycle and its transmission at mitosis.

Unlike classical mRNAs which are exported to the cytoplasm or to other ribonuclear particles and which are characterized by their high mobility [28], the Xist RNAs remain localized within the nucleus and tightly associated with the inactive chromatin. Experiments aimed at dissociating the Xist containing chromatin fraction have indicated that Xist RNAs is bound to the nuclear matrix and suggested an association of Xist transcripts with nuclear matrix attachment proteins [29]. The accumulation of scaffold associated factor A (SAF-A) on the inactive X may indicate that this factor ensures the link between Xist and the nuclear matrix [30]. Restriction of Xist RNAs to the nucleus may also involve a specialized nuclear compartmentalization process.

The Xist transcript represents the canonical example illustrative of the ability of some ncRNAs to ‘travel’ long distances in order to induce chromatin modifications. It is interesting to note that despite its long range effect, the Xist RNA propagates itself only along the chromosome from which it originates. The molecular mechanisms responsible for this cis-spreading and the consequent chromatin refolding remain poorly understood. Xist RNA propagation is thought to involve relay or entry sites interspersed along the chromosome. These specialized sequences may work as nucleation centres which concentrate chromatin remodelling complexes and facilitate the propagation of the silent chromatin structure to surrounding sequences (Fig. 2A). One such nucleation centre has been proposed to correspond to a 150-kb region lying just 5’ to the Xist gene which is constitutively enriched in H3K9 methylation [31] (Rougeulle et al., in press) (Fig. 3).

The inactive X chromatin is also enriched in the histone variant macroH2A, which accumulates within a nuclear structure known as the macrochromatin body. Xist RNA is necessary for the localization of macroH2A to the inactive X [32]. Due to its histone nature, macroH2A has been proposed to be involved in linking the X-chromatin to Xist RNA. Since, in the ES cell model, macroH2A only becomes associated with the inactive X, a considerable time after the onset of XCI, it is however more likely to be involved in the maintenance of the transcriptionally silent chromatin structure.

Site-specific transgenes expressing different forms of Xist RNA have allowed the function of specific parts of the Xist transcript to be analysed [33]. Interestingly, a repeat sequence located at the 5’ extremity of the Xist RNA which is able to adopt a double hairpin conformation seems to be responsible for the silencing function of Xist. In the absence of this motif, Xist decorates the X-chromosome without however inducing transcriptional repression. In contrast, the coating function appears to be mediated by many different and redundant sequences within the transcript acting in a cooperative manner.

In Drosophila, unlike the situation pertaining in mammals, dosage compensation is achieved by a 2-fold upregulation of transcription of genes on the single X-chromosome present in the males. Intriguingly, however, the fly dosage compensation system also involves two ncRNAs: roX1 and roX2 (RNA on the X) which show features similar to Xist. roX1 and roX2 are members of the dosage compensation complex (DCC), which also contains the male specific lethal (MSL) proteins. An intact DCC complex is required for hyper-transcription of genes on the X-chromosome. A translation block of any one of these subunits results in the failure of the DCC assembly in females (for review see [34]). MSL proteins include chromodomain proteins which can interact with RNA in vitro and an histone acetyl transferase responsible for the decondensed chromatin structure which characterizes the dosage compensated Drosophila X-chromosome. In genetic backgrounds where some of the MSL proteins have been mutated, a partial DCC can still be detected associated with approximately 35 primary “chromatin entry sites” distributed along the X [35]. These “entry sites” have been shown to act as initial docking sites for the DCC assembly from where the complex spreads in cis to hundreds of other sites which ensure the propagation of a decondensed chromatin structure along the entire X-chromosome (Fig. 2A).

Two of these examples serve to emphasize the capacity of some ncRNAs to spread in cis over long distances (it is, however, noteworthy that roXs RNAs can also act in trans when inserted at an ectopic location). Interestingly, the RNA molecules described in these examples seem to be capable of multiple interactions with chromatin modifying enzymes; these interactions mediating the chromatin organization at the chromosomal level. The recent identification of the involvement of an RNA component in the establishment of histone modification patterns at mammalian pericentric heterochromatin may point to a more general implication of RNA molecules in high-order chromatin structures [36].

3. Antisense ncRNAs and transcriptional repression

Xist RNA is the major effector responding to the initiation signal(s) involved in XCI. Intriguingly, Xist expression itself is controlled by another non-coding transcription, antisense to Xist, named Tsix (Fig. 3). Tsix transcription is mainly initiated 12 kb downstream of Xist and gives rise to approximately equivalent amount of primary and spliced transcripts which span a 40-kb region and overlap the entire Xist locus. Like Xist, Tsix is expressed on all Xs prior to the onset of XCI. When XCI initiates, Xist RNA up-regulation on the future inactive X-chromosome is accompanied by transcriptional repression of Tsix, whereas on the active X Tsix expression persists [37].

Genomic imprinting, like XCI, leads to monoallelic expression of target genes. Non-coding antisense transcription is frequent as already mentioned at imprinted loci and shows a mutually exclusive expression pattern with its sense counterpart. Antisense transcription at imprinted loci can extend over several hundred kilobases (Air, Antisense transcription at the Igf2r locus, 108 kb; Ube3a-as, antisense at the Angelman Syndrome/Prader-Willi Syndrome locus (AS/PWS locus), 450 kb or more, Fig. 1B) The extended nature of the
transcripts may account for and mediate the long-range cis-effects associated with the antisense transcription. Similarities between the regulation at the Xic and imprinted regions may indicate that non-coding antisense transcription at both exploit the same underlying molecular mechanism(s). Truncation of both the Air and the Tsix antisense transcripts by insertion of premature polyadenylation signals [38,39] induces the reactivation of the sense transcription at both loci con-
comitant, respectively, with the loss of imprinted gene expression and distortion of random XCI. These results suggest a direct implication of antisense transcription in monoallelic repression of overlapping gene function. However, it is important to appreciate that these experiments do not allow discrimination between the act of antisense transcription per se as opposed to the direct implication of antisense RNA molecules.
Although recent studies have identified many NATs, the molecular mechanisms underlying NAT-mediated gene repression are still unknown. Several different models based on results obtained in eukaryotic organisms have been proposed. Transcription-dependent mechanisms hypothesize that the repressive function originates in the movement of the RNA polymerase machinery in the opposite direction to sense transcription leading to topological constraints and/or molecular collisions involving sense transcription (Fig. 2B). This hypothesis is favoured by the observation that antisense transcripts tend to overlap the 5′ region of their sense genes which regroup essential elements such as the promoter, translation initiation sites and enhancers, and would be expected therefore to be particularly vulnerable to antisense transcription mediated disruption. However, this type of mechanism cannot fully explain the repressive effect at the Air Igf2r locus as genes outside of the region overlapping Air are also transcriptionally repressed. Since, prior to XCI, Xist and Tsix are co-expressed on both X-chromosomes, for a transcriptional interference model to be relevant, it would be necessary to postulate that the Xist and Tsix genes are transcribed alternately.

Another class of model proposes a direct role for the antisense ncRNA molecules (Fig. 2B) probably via a pairing between sense and antisense transcripts leading to the formation of dsRNAs. The formation of dsRNAs may either, through a destabilization process, induce the degradation of the sense/antisense complex, or alternatively mask access to functional domains of either or both transcripts by trans-acting factors. This RNA masking might interfere with any of the multiple steps involved in post-transcriptional RNA-processing including splicing, polyadenylation, transport and translation. In the case of Xist/Tsix, it is interesting to note that the absence of Tsix is associated with the dispersion of Xist RNAs outside of their transcription site even prior to the onset of XCI [40]. This observation suggests an additional role for antisense transcripts in anchoring sense RNAs at their site of transcription or at sites of anchorage. The significant molar excess of Tsix RNAs over Xist transcripts [41] may allow the trapping of all the Xist RNAs produced and in a certain manner, obstruct the access of the transcriptional machinery to the Xist promoter, leading to partial repression of Xist expression (Fig. 2B).

The very widespread use of RNAi-based mechanisms in different biological systems and recent evidence demonstrating that at least in some species, a part of the RNAi pathway may occur in the nuclear compartment [42] suggest a role for this process in antisense-mediated gene repression. According to this view, dsRNAs would be cleaved into short interfering RNAs (siRNAs) by Dicer or other RNaseIII family members, which would mediate the silencing of the surrounding genes presenting a minimal sequence homology (for review [43]). In the PTGS model, inhibition of the target genes results from degradation of the RNA products by siRNA. In the transcriptional gene silencing (TGS) situation, silencing is mediated by DNA methylation and chromatin changes. Links between RNAi and chromatin states have recently been established in fission yeast, where the RNAi machinery can apparently direct H3K9 methylation and repress H3K4 methylation at centromeric repeats thereby inducing heterochromatic silencing [42]. At outermost region (otr) repeats, complementary sense and antisense transcriptions induce the formation of siRNAs, which are thought to guide histone modifications and the recruitment of heterochromatin proteins. A provocative correlation between H3K4 methylation and Tsix expression has recently been observed within the Xist gene, which may suggest the involvement of an RNAi pathway in XCI. However, neither dsRNAs nor siRNAs corresponding to the Xist/Tsix overlapping region have as yet been described despite intensive investigation. The size of the region and the nature of the transcripts obviously complicate such analysis. It is, however, noteworthy that the XCI process would necessitate the involvement of a cis-limited RNAi pathway.

4. Non-coding transcriptions, open chromatin structures and trans interactions

Within known regulatory complexes, additional transcription units involving transcription from intergenic loci are increasingly being described. Three examples of this phenomenon from amongst many (for review [13]) will be discussed on here. Within the Xic, two novel sites of intergenic transcription have been reported so far. The first, known as the “B region”, is located approximately 50 kb 3′ and upstream to Xist (Fig. 3). Transcriptional activity is initiated from this region in both orientations [18]. The second, the X-inactivation intergenic transcription element (Xite) locus, lies 30 kb 3′ and downstream to Xist and is associated with a series of DNaseI hypersensitive sites. Both the transcriptional activity and DNaseI sensitivity of the Xite locus are developmentally regulated with an expression profile that parallels that of the Tsix gene during

Fig. 3. Map of the mouse Xic. This simplified map shows the 200 kb of the Xic around the Xist locus and the locations of known non-coding transcription units within this region. The extent of the H3K9 methylation hotspot, which may act as a nucleation centre allowing the spreading of the X-inactivation signal, is also shown.
XCI [44]. This observation has led the authors to propose that Xite might act as an enhancer or as a LCR modulating Tsix expression.

Such intergenic transcription units produce ncRNAs of variable size. The role of these molecules has not been addressed up until now as it has been considered that their function is likely linked to the use of transcription-dependent mechanisms rather than being RNA-dependent per se. This assessment has been confounded by the discovery of an association between the C-terminal domain of the elongating RNA polymerase II holoenzyme and histone modifying enzymes marking active chromatin domains such as histone acetyl transferases [45,46] and H3K4 histone methyl transferases [47,48]. This link between the transcriptional machinery and chromatin remodelling enzymes has led to the seductive piggybacking hypothesis [49,50]. According to this model, the movement of the RNA polymerase II elongation complex along the chromatin fibre would allow the deposition of histone modifications (Fig. 2C). Seen from this perspective, non-coding transcription would have as its ultimate function the definition of domains of open chromatin necessary for the facilitated binding of trans-acting factors and other trans-interactions with other genomic elements. Transcription at the Xite locus could create docking sites accessible for transcription factors and would favour interactions between these factors and cis-regulatory elements associated with target genes thereby allowing transcriptional activation.

Tsix antisense transcription itself may also participate in the definition of an open chromatin domain, as the loss of Tsix antisense transcription is associated with a loss of H3K4 methylation within the Xist gene [51]. This may suggest that the Tsix non-coding antisense transcription intervenes both in the repression of Xist expression and in the formation of an open chromatin domain which is potentially involved in other parts of the XCI process. XCI by its nature must involve trans-interactions between X-chromosomes as well as interactions with autosomal factors that could require the establishment of an open chromatin structure around Xist to expose target sequences. More generally, one might expect that non-coding intergenic transcription will, through chromatin remodelling, be involved in cis and trans communications between elements distributed over large genomic domains such as the Xic.

The correlation between intergenic transcription and regulatory activity is also exemplified by regulation at the Bithorax complex (BX-C) during Drosophila embryo development (Fig. 1B). The BX-C complex, which extends over 300 kb, includes three coding genes: the homeotic genes Ultrabithorax, Abdominal-A (abdA) and Abdominal-B (AbdB) [52]. The sequential expression of these genes along the antero-posterior embryo axis determines the fate of each parasegment of the embryo [53]. Regulation of the abdA and AbdB genes is under the control of the infra-abdominal (iab) region. This 100-kb cis-regulatory region has been genetically divided into numerous iabs subdomains (iab 2–8) containing both enhancers and silencers. The iabs are transcribed following the same collinearity rule as homeotic genes [54]. Alteration of transcription in one iab subdomain induces a homeotic transformation of the more posterior segment under its control, suggesting that intergenic transcription plays a crucially important role in iabs activity [55].

A final example concerns the human β-globin locus (Fig. 1B). This locus consists of five genes tightly regulated during the erythropoietic development (for review see [56]). The regulation of the β-globin locus is governed by an LCR, located 8 kb upstream of the first β-globin gene and which contains five DNaseI hypersensitive sites. This LCR shows constitutive intergenic transcription activity, which is thought to intervene in the sequential gene activation of β-globin genes by helping in the establishment of three functional domains. The extent of these domains is precisely delineated by intergenic transcription occurring at the β-globin locus downstream of the LCR and correlates exactly with the activation patterns of the β-globin genes themselves. These domains are characterized by chromatin modifications, which also correlate with the intergenic expression state [57].

Many of these results suggest that the mechanism of repression mediated by ncRNAs may vary with the regulatory system. Whilst antisense transcription seems to be especially frequently employed in the establishment of allele-specific expression of target genes and many of our notions concerning ncRNAs have been derived from the study of such systems, the analysis of antisense transcripts at additional loci may well reveal other modes of functioning and regulatory involvement.

The discovery that ncRNAs and non-coding transcription play a role in varied and highly disparate regulatory systems and exploit an extended variety of mechanisms suggests that ncRNAs are likely implicated and integral to the overall molecular architecture of organisms (for review see [58]). The expanding numbers of new, functional ncRNAs together with the observation that regulatory RNA molecules can be produced from the introns of coding genes suggests a role for ncRNAs as signalling molecules that can be both easily produced and destroyed at a smaller energetic cost than can protein signalling molecules which require both protein synthesis and proteolysis. In this view, the transcriptional background provided by “junk” DNA may not only participate in the maintenance of genome wide low-level transcriptional activity but also be required for efficient and integrated cellular function. The baroque hypothesis of a primordial “RNA world” clearly deserves to be revisited. We may also be obliged to reevaluate our notions of what constitutes ‘junk’ DNA.

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