

### **PRIMER**

# CncRNAs: RNAs with both coding and non-coding roles in development

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#### **ABSTRACT**

RNAs are known to regulate diverse biological processes, either as protein-encoding molecules or as non-coding RNAs. However, a third class that comprises RNAs endowed with both protein coding and non-coding functions has recently emerged. Such bi-functional 'coding and non-coding RNAs' (cncRNAs) have been shown to play important roles in distinct developmental processes in plants and animals. Here, we discuss key examples of cncRNAs and review their roles, regulation and mechanisms of action during development.

KEY WORDS: Bifunctional RNA, CncRNAs, Coding RNA, Development, Non-coding RNA, Regulatory RNA

#### Introduction

Ribonucleic acids (RNAs) serve important roles, most notably as intermediaries in the flow of genetic information from DNA to proteins. They can be found in various forms, for example as protein-coding messenger RNAs, as recruiters and machines for protein synthesis (e.g. transfer RNAs, ribosomal RNAs), as modifiers of ribosomal RNAs, and as regulators of RNA splicing, RNA stability and protein synthesis (e.g. small nuclear RNAs, small nucleolar RNAs and microRNAs). Other small RNAs function in epigenetic regulation and post-transcriptional gene silencing, and protect the genome from transposons (e.g. piwi-interacting RNAs). In addition to these well-studied groups of RNAs, transcriptomic analyses in a variety of organisms have identified hundreds of other RNAs (e.g. long non-coding RNAs) that are thought to function in numerous cellular and developmental processes (Bushati and Cohen, 2007; Houwing et al., 2007; Mercer et al., 2009; Pauli et al., 2011; Rinn and Chang, 2012; Ulitsky and Bartel, 2013; Weick and Miska, 2014; Hezroni et al., 2015).

Until recently, most RNAs were presumed to be exclusively protein coding or non-coding. However, studies of many bacteria, animals and plants have revealed an unusual group of RNAs that have both protein coding and non-coding roles. These RNAs have been referred to as 'dual function' or 'bi-functional' RNAs (Dinger et al., 2008; Ulveling et al., 2011b); we refer to them hereafter as 'coding and non-coding RNAs' (cncRNAs) (Kumari and Sampath, 2015). The identification of such cncRNAs raises several interesting questions and poses a challenge to RNA classification. Here, we summarize the known features of a few exemplary cncRNAs and discuss their roles and regulation during plant and animal development.

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#### cncRNAs in animal development

A number of cncRNAs that function in differentiation and development in animals have been identified (Table 1). The dual functionality of these RNAs was often revealed serendipitously by comparing RNA-null and protein-null mutant alleles, by disrupting RNA elements, by overexpression assays and via antisense-mediated depletion.

#### Oskar

The *Drosophila melanogaster* protein Oskar (Osk) plays essential roles during germline and abdominal segment formation in the developing fly embryo (Lehmann and Nusslein-Volhard, 1986; Ephrussi et al., 1991; Kim-Ha et al., 1991). Osk protein is produced at the posterior pole of the *Drosophila* oocyte, where it recruits the germline specific RNA helicase Vasa in the process of germ plasm assembly; Osk has also been shown to bind to germline RNAs and Vasa in vivo and in vitro (Breitwieser et al., 1996; Jeske et al., 2015; Yang et al., 2015). Interestingly, osk RNA has roles independent of Osk protein in early oocytes. This non-coding activity of osk RNA was initially deduced from the distinct behaviors of classical EMSinduced osk nonsense mutants that produce osk mRNA but no detectable Osk protein (Lehmann and Nusslein-Volhard, 1986) and transposon-induced osk RNA-null mutants (Jenny et al., 2006). osk is a maternal effect gene, expression of which is required in the female germline for proper development. However, lack of the mRNA and lack of the protein lead to two very different outcomes: in the absence of Osk protein, the progeny embryo fails to form germ cells and abdominal segments (Ephrussi et al., 1991; Kim-Ha et al., 1991), whereas the complete absence of osk mRNA causes an arrest in oogenesis, and no eggs are produced (Jenny et al., 2006). It was further shown that transgenes expressing mutant osk RNAs that cannot make functional Osk protein, as well as those harboring only the osk 3' untranslated region (3'UTR), can overcome the early oogenesis defects of osk RNA-null mutants. This suggested that osk RNA harbors a non-coding activity in its 3'UTR that functions during oogenesis (Jenny et al., 2006).

A more recent study showed, remarkably, that the expression of a transgenic 191-nucleotide segment of the *osk* 3'UTR is sufficient to rescue the oogenesis arrest of *osk* RNA null oocytes, as long as the RNA is fused to a stem-loop structure that promotes its dynein-dependent import into the oocyte (Kanke et al., 2015). It was further shown that mutations in the *osk* 3'UTR that abolish binding of the translational regulator Bruno (Bru; Aret – FlyBase) to Bru response elements (BREs) in *osk* also affect egg laying (Kanke et al., 2015). The comparison of egg-laying by females lacking endogenous *osk* RNA with those expressing *osk* transgenes that disrupt some or all the BREs showed that the mutation of any BRE reduces egg-laying to some extent, with the strongest effect observed when all BREs are lost. The reduction in egg-laying appears to be due to disruption of the non-coding function of *osk* RNA (Kanke et al., 2015).

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Table 1. Coding and non-coding roles of known cncRNAs

| Organism   | Gene                                   | Peptide/protein function   | Non-coding RNA function   |
|--|--|--|---|
| Drosophila melanogaster  | oskar                                  | Germ-line RNA and Vasa helicase-binding protein; required for germ-line and abdominal specification in embryos | Oogenesis (egg production)  |
| Zebrafish (Danio rerio)  | squint (nodal-<br>related 1)           | Secreted signaling morphogen; required for<br>mesendoderm specification  | Dorsal axis formation   |
| Xenopus laevis   | vegt                                   | T-box-containing transcription factor;<br>mesoderm and endoderm formation                                      | Germinal granule formation; cytoskeletal anchoring  |
| Rattus norvegicus  | Ube3a                                  | Ubiquitin ligase; protein turnover; synapse<br>development; plasticity, learning and memory                    | Dendritic growth and spine maturation in hippocampal neurons                              |
| Mouse (Mus musculus)   | Steroid receptor activator (Sra; Sra1) | Regulation of estrogen hormone; spliceosome complex protein  | Transcriptional co-activator of steroid<br>hormone receptors; myogenic<br>differentiation |
| Human  | Insulin receptor substrate 1 (IRS1)    | Co-factor of insulin and insulin-like receptor; effector of insulin signaling                                  | Regulation of <i>RB</i> mRNA and myoblast proliferation                                   |
| Medicago trunculata, Glycine<br>max, Arabidopsis thaliana,<br>Oryza sativa | enod40                                 | Regulation of sugar metabolism; root and cotyledon growth  | Root nodule formation   |
| Medicago trunculata  | miRNA171b                              | miPEP that upregulates pri-miRNA171b   | Repression of Scarecrow-like<br>transcription factors and auxiliary<br>meristems          |
| Human  | p53                                    | Tumor suppressor protein; cell cycle regulation and growth control   | Regulator of MDM2   |

The non-coding activity of *osk* could be mediated either by sequestering an oogenesis inhibitory factor, or by a scaffolding function for a ribonucleoprotein (RNP) that assembles on *osk* RNA and promotes oogenesis. Evidence that the *osk* 3'UTR acts via sequestration was provided by mutations affecting Bru, which binds to Osk; mis-sense mutations in *bru* that reduce Bru levels suppress the *osk* null mutant phenotype, and this effect is even stronger with *bru* nonsense mutant alleles. However, mutations in regions of *osk* mRNA that are not involved in Bru binding also affect egg-laying, indicating that additional unknown factors contribute to the noncoding functions of *osk* (Kanke et al., 2015).

#### Squint

The zebrafish protein Squint (Sqt; Nodal-related 1 – Zebrafish Information Network) is a secreted signaling morphogen that is essential for formation of the embryonic organizer and for specification of mesendoderm during gastrulation (Erter et al., 1998; Feldman et al., 1998; Rebagliati et al., 1998; Chen and Schier, 2001). However, maternal sqt RNA has an earlier non-coding function, independent of Sqt/Nodal signaling, during the formation of the zebrafish dorsal axis (Gore et al., 2005, 2007; Bennett et al., 2007; Lim et al., 2012). This non-coding activity was identified by overexpression and antisense knockdown studies: overexpression of mutant zebrafish sqt RNA that is incapable of encoding functional Sqt/Nodal protein expands the expression domains of dorsal progenitor genes such as goosecoid and chordin. Furthermore, the dorsal-inducing activity of sqt RNA does not require the Sqt-coding exons, and the 3'UTR of sqt RNA is sufficient for its dorsalizing activity.

It was also found that targeting maternal sqt RNA with antisense morpholinos that reproduce zygotic sqt mutant phenotypes causes mislocalization of sqt RNA and the complete loss of embryonic dorsal structures. This is in contrast to insertion mutations in the sqt locus that disrupt Sqt protein coding sequences but do not affect sqt RNA (Feldman et al., 1998; Feldman and Stemple, 2001; Aoki et al., 2002; Amsterdam et al., 2004; Gore et al., 2005; Lim et al., 2012). The precise molecular mechanism by which non-coding sqt RNA elicits its role is not known. Systematic analysis of mutants

lacking sqt RNA can help address this question (Lim et al., 2013). Interestingly, the UTR-mediated non-coding activity of sqt RNA is independent of microRNA (miRNA) target sequences and the function of the miRNA processing gene dicer (dicer1 – Zebrafish Information Network) (Lim et al., 2012). Therefore, binding to miRNA negative regulators of dorsal identity is unlikely to be the mode of action of non-coding sqt RNA. However, a requirement for the Wnt/ $\beta$ -Catenin pathway raises the possibility that sqt RNA might act as a scaffold for a factor that promotes nuclear accumulation of  $\beta$ -Catenin in dorsal progenitors. Alternatively, non-coding sqt RNA might sequester a negative regulator of dorsal identity.

## VegT

VegT is a T-box family transcription factor that functions in mesoderm and endoderm formation during gastrulation in *Xenopus laevis* embryos (Zhang et al., 1998; Heasman et al., 2001; Kloc et al., 2005). However, in *Xenopus laevis* oocytes, *vegt* RNA plays an additional role in the organization of cytokeratin filaments and germinal granules. Accordingly, depletion of *vegt* RNA in *Xenopus* oocytes by phosphorothioate antisense oligonucleotides results in disruption of the cytokeratin network, mislocalization of maternal RNAs, and blocked formation of germinal granules (Heasman et al., 2001; Kloc et al., 2005). Exogenously provided *vegt* RNA was found to reconstitute and rescue the disrupted cytokeratin network (Kloc et al., 2005). Based on these lines of evidence, *vegt* RNA is thought to be a structural matrix that stabilizes cytokeratin filaments and anchors vegetal RNAs in frog eggs.

## Steroid receptor activator RNA

The mammalian steroid receptor RNA activator (SRA; also known as SRA1) promotes myogenic differentiation and the conversion of non-muscle cells to myocytes by co-activation of the myogenic differentiation factor MYOD (also known as MYOD1). In addition to MYOD, *SRA* transcripts are believed to co-activate numerous nuclear receptors, and potentially regulate the proliferation and differentiation of a variety of cell types. *SRA* RNA has been found associated with the PRC2 Polycomb group and TrxG trithorax

transcriptional repressor complexes, and it directly interacts with the stem cell pluripotency factor NANOG (Wongtrakoongate et al., 2015). By contrast, SRA protein (SRAP) prevents *SRA* RNA-mediated co-activation and differentiation (Lanz et al., 1999; Hube et al., 2011), and is thought to act as a trans-activator of steroid hormone receptors in mammalian cell culture assays.

The activity of *SRA* RNA as a trans-activator of steroid hormone receptors and as a co-activator of MYOD was identified by studying *SRA* transcripts lacking a methionine initiation codon. Initial attempts to identify *SRA* transcripts with an extended 5' sequence were unsuccessful, leading to the hypothesis that SRA functions as a non-coding RNA in this context. Subsequently, mutations in *SRA* revealed that its co-activator function resides in the transcript (Lanz et al., 2002). It was also shown that introducing multiple stop codons in *SRA* RNA did not abolish steroid receptor co-activation, and RNA activity was detected even in the presence of the translation inhibitor cycloheximide. Together, these findings suggested that *SRA* RNA has functions independent of SRAP (Lanz et al., 1999).

A recent study reporting the crystal structure of human SRAP suggests that the protein does not harbor a predicted RNA-recognition motif (RRM) but rather resembles the spliceosome complex protein PRP18 (also known as PRPF18) (McKay et al., 2014). Biochemical binding assays performed *in vitro* did not find a specific interaction between SRAP and *SRA*, nor was a specific response on estrogen receptor targets observed in cell culture assays. These observations have led to the proposal of an alternative model wherein SRAP is thought to stabilize intermolecular interactions within a nuclear splicing complex, rather than directly binding and regulating *SRA* RNA (McKay et al., 2014). Genetic analysis of the *sra* locus with mutations that disrupt specific *SRA* RNA versus SRAP domains, which could resolve the roles and mechanisms by which the RNA and protein function during development, differentiation and homeostasis, is currently lacking.

## p53

The tumor suppressor protein p53 (TP53) plays crucial roles in cell cycle regulation and in preventing mutations arising from DNA damage in the genome (Lane and Crawford, 1979). However, in addition to the well-known roles of p53 in protecting the genome, p53 mRNA was found to regulate the ubiquitin ligase MDM2, which is a negative regulator of p53 (Candeias et al., 2008). p53 mRNA directly interacts with the N-terminus of MDM2 to prevent its E3 ubiquitin ligase activity and thereby controls MDM2mediated regulation of p53. Interfering with the p53 mRNA-MDM2 interaction following DNA damage prevents p53 stabilization and activation (Gajjar et al., 2012). Thus, p53 mRNA-MDM2 interactions are key for the genotoxic stress response. Interestingly, the sequences of p53 mRNA that interact with the N-terminus of MDM2 protein also encode the amino acids in TP53 that interact with and are poly-ubiquitylated by the MDM2 RING domain (Naski et al., 2009). Silent point mutations in this region of p53 weaken its interactions with MDM2, and reduce TP53 activity. This suggests that structural elements in p53 mRNA might harbor its non-coding activity.

#### IRS1

Insulin receptor substrate 1 (IRS1) is a major substrate and cytoplasmic docking protein for the insulin receptor and insulin-like growth factor receptor. IRS1 is thought to be an effector of insulin signaling, with roles in cell growth and proliferation. Deletion of *Irs1* by conventional knockout strategies in mice led to

growth retardation and compensated insulin resistance (Kido et al., 2000). IRS1 levels are generally low or absent in differentiating cells, and elevated IRS1 levels have been associated with cancers in mice and humans. Interestingly, a recent study found that Irs1 mRNA has a function in myoblasts that is independent of IRS1 protein (Nagano et al., 2015). The 5'UTR of Irs1 mRNA harbors sequences complementary to RNA encoding the cell cycle regulator retinoblastoma (RB; RB1). Overexpression of the 5'UTR region of Irs1 led to reduced Rb mRNA expression, whereas knockdown of Irs1 mRNA harboring the 5'UTR complementarity region led to increased Rb mRNA levels and enhanced myoblast differentiation. These findings suggest that Irs1 RNA has a novel role as a regulatory RNA that is independent of IRS1 protein (Nagano et al., 2015). It is not known whether *Irs1* mRNA regulation is restricted to myoblasts, and what controls Irs1 RNA versus IRS1 protein functions is also unknown.

#### **UBE3A**

Finally, a recent study reports that transcripts encoding the E3 ubiquitin ligase UBE3A have a non-coding role in dendrite growth and spine maturation in hippocampal neurons; by contrast, UBE3A protein plays crucial roles in activity-dependent synapse development, plasticity, learning and memory (Sun et al., 2015; Valluy et al., 2015). This role was discovered using antisense short hairpin RNAs (shRNAs) that target Ube3a 3'UTR variants in rat hippocampal neurons: the neurons showed significantly increased dendrite growth and complexity upon knockdown of the non-translated and intron-retaining *Ube3a1* RNA, whereas shRNAs targeting the spliced coding Ube3a2/3 isoforms reduced dendrite complexity (Valluy et al., 2015). Antisense oligonucleotides can have off-target effects or, alternatively, they can uncover novel mechanisms not identified by protein-disrupting mutants. Therefore, it is crucial that the non-coding activities of cncRNAs identified by antisense approaches (e.g. shRNA, antisense morpholinos, phosphorothioate oligonculeotides) are independently validated by assays such as overexpression of nontranslatable RNA, and by analysis of mutations that specifically disrupt the RNA (Lim et al., 2012, 2013; Kok et al., 2015; Rossi et al., 2015).

#### **Plant cncRNAs**

A number of cncRNAs have also been identified in plants (Table 1). Transcripts of the gene *early nodulin 40 (enod40)* in the legume *Medicago trunculata* contain two short open reading frames (ORFs), the products of which are required for cortical cell divisions in root cells (Yang et al., 1993; Crespi et al., 1994). A region of RNA secondary structure separates the two *enod40* ORFs. This RNA segment is essential for *enod40* activity and has a noncoding role in root nodule formation (Girard et al., 2003; Campalans et al., 2004). It was also shown that, in alfalfa, *enod40* RNA is essential for a growth response in the root cortex (Sousa et al., 2001).

In soybean plants, ENOD40 peptides have been shown to regulate the turnover of the enzyme sucrose synthase (SUC1), which functions in sugar metabolism in roots and cotyledons. By using a combination of RNA structure prediction, comparison and structure probing, various regions of soybean *enod40* RNA were identified to be key for root nodule formation. Of these, five domains are conserved amongst leguminous plants and are presumed to be required for the non-coding activity of *enod40* RNA (Girard et al., 2003). Indeed, the deletion of an inter-ORF RNA region with predicted structure resulted in reduced activity of alfalfa *enod40* 

without affecting the production of ENOD40 peptides (Sousa et al., 2001).

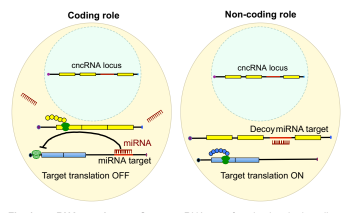
Analyses in *Arabidopsis thaliana* and rice have identified RNAs similar to *enod40* suggesting that cncRNAs exist in these plants as well (Kouchi et al., 1999). Furthermore, at least 50 miRNAs in the *Arabidopsis* transcriptome are predicted to encode short peptides (microRNA-encoded peptides; miPEPs) that appear to regulate the transcription of primary transcript (pri-)miRNA. For example, precursor (pre-)miRNA for *Medicago trunculata* miR171b encodes a short peptide expression of which leads to transcriptional upregulation of the corresponding pri-miRNA, which in turn controls target genes involved in root development (Lauressergues et al., 2015). Interestingly, all the identified miPEPs were found to be conserved across flowering plants and are associated with ancient miRNA families. It is not known if miRNA-encoded peptides are present in animals.

#### cncRNA functions and mechanisms of action

The precise function and molecular mechanism of action is known for only a few cncRNAs. Nonetheless, it is emerging that all the mechanisms deployed by conventional non-coding RNAs are also represented amongst this hybrid class of RNAs.

#### Base pairing and roles as decoy and regulatory RNAs

Base pairing is a fundamental property of nucleic acids that is essential for processes such as codon-anticodon recognition during protein synthesis and microRNA recognition of target RNA sequences. Nucleotide complementarity also forms the basis for RNAs that function as 'molecular sponges', 'decoys' or 'target mimics', such as competing endogenous RNAs (ceRNAs) that share miRNA recognition sequences. ceRNAs bind to complementary sequences in miRNAs and prevent interactions between miRNAs and their bona fide targets (Franco-Zorrilla et al., 2007). For instance, the pseudogene *PTENP1* harbors sequences in its 3'UTR that are complementary to miRNAs that target and repress PTEN mRNA (Poliseno et al., 2010). Indeed, cncRNAs can also function by base pairing with target RNAs and behave as decoys (Fig. 1). For example, non-coding *Ube3a1* RNA retains an intron and is thought to act as a decoy or molecular sponge for miRNA-134 (MIR134) that would otherwise target the spliced Ube3a2/3



**Fig. 1. cncRNAs** as **decoys.** Some cncRNAs can function in a both coding role (left), being translated into protein (yellow chain) and having no effect on target genes, and a non-coding manner (right), by base pairing with and hence acting as decoys for small regulatory RNAs (e.g. miRNAs) that repress target translation. In the example shown, intron retention leads to the presence of a decoy miRNA target site in the cncRNA that can bind to an miRNA (red comb) and allow translation of the target of this miRNA (blue chain). Ribosomes are shown in green.

transcripts and downregulate UBE3A ubiquitin ligase protein expression in dendrites (Valluy et al., 2015).

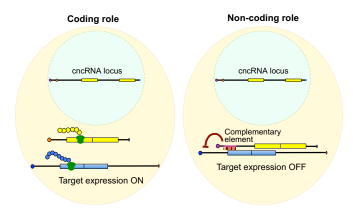
Some cncRNAs appear to target and regulate other mRNA sequences directly (Fig. 2). For example, the 5'UTR of mammalian *Irs1* RNA can base pair with *Rb* mRNA. The overexpression of *Irs1* RNA, which contains a sequence element complementary to *Rb* mRNA, reduces RB levels by a mechanism independent of DICER or UPF1 (i.e. independently of miRNAs or nonsense-mediated decay), and suppresses the differentiation of cultured skeletal muscle cells (Nagano et al., 2015). Interestingly, some bacterial small regulatory RNAs (sRNAs) also function in a similar manner. For example, *Escherichia coli SgrS* RNA regulates the glucosephosphate stress response by base pairing with and blocking the translation of *PtsG* mRNA, which encodes a sugar phosphate transporter. *SgrS* is a cncRNA as it also encodes a short 43 amino acid peptide, SgrT, which inhibits the activity of the PtsG transporter protein (Wadler and Vanderpool, 2007; Rice and Vanderpool, 2011).

#### Structural roles, sequestration and scaffolding

Structural features in RNAs play crucial roles in their activity. RNA structural elements can either sequester or bind and deliver protein complexes (Figs 3 and 4). SRA RNA co-activation of MYOD, for example, is mediated via interactions between SRA RNA and the p68/p72 RNA helicases (Caretti et al., 2006). A sequestration and scaffolding function has also been proposed for osk 3'UTR sequences during early oogenesis, via their binding to Bru and some unknown factors (Kanke et al., 2015). In Xenopus eggs, vegt RNA forms aggregates that colocalize with cytokeratin filaments, and the depletion of vegt destabilizes the cytokeratin network and disrupts the anchoring of vegetal RNAs (Heasman et al., 2001; Kloc et al., 2005), suggesting that vegt RNA also plays a scaffolding or sequestering role. Finally, it was shown that the RNA sequence separating the two ORFs within *enod40* RNA harbors a region of highly stable secondary structure that binds to the RNA-binding protein MtRBP1, which is presumed to be a translational regulator (Crespi et al., 1994; Sousa et al., 2001; Campalans et al., 2004). Structural elements in RNAs can also act as sensors for environmental stimuli, as observed in some bacterial RNAs: these RNAs function as inactive nascent transcripts with secondary structures that are released upon binding of the RNA to ligands or metabolites, or upon increase in temperature, leading to activation of downstream gene expression. One such example is the S-adenosylmethionone-sensing S-box riboswitch in *Bacillus* subtilis (Henkin, 2008; Gottesman and Storz, 2011). Structural RNA sensors could facilitate rapid changes in gene expression in response to external signals during developmental transitions (Fig. 5).

## Feedback regulation

Some cncRNAs mediate feedback regulation of the same pathway or process in which their protein product functions. This type of feedback is exemplified by the p53-MDM2 pathway (Candeias et al., 2008). A region of p53 mRNA binds to and regulates the E3 ubiquitin ligase and p53 tumor protein regulator MDM2, which inhibits p53 activity by controlling p53 translation, polyubiquitylation and degradation. The binding of p53 mRNA to MDM2 leads to accumulation of MDM2 at polysomes, stimulation of p53 synthesis, and inhibition of the E3 ligase activity of MDM2. Accordingly, mutations in p53 that reduce the affinity of p53 mRNA for MDM2 enhance the suppression of p53 activity by MDM2. Thus, p53 mRNA acts as a feedback switch that controls MDM2-mediated regulation of p53 by directly interacting with MDM2 (Candeias et al., 2008). Such regulation is also seen in the context of



**Fig. 2. cncRNAs as regulatory RNAs**. Complementary sequence elements in cncRNAs can bind directly to other mRNAs and control their expression. In the example shown, the coding cncRNA is transcribed from one transcription start site (orange circle) and translated (left; yellow chain), whereas the non-coding cncRNA (right) is generated from an alternative transcription start site (magenta circle) such that the 5'UTR now includes a region of complementarity (red comb) to the target mRNA. Ribosomes are shown in green.

MYOD activation: *SRA* ncRNA acts as a co-activator of MYOD during myogenic differentiation, whereas the protein SRAP has an inhibitory effect on *SRA* RNA and hence MYOD co-activation. SRAP is thought to exert its inhibitory effect by binding to *SRA* RNA (Hube et al., 2011). In both of these examples, feedback regulation is achieved by the cncRNA binding to a protein that functions in the same developmental or cellular process as does the cncRNA.

#### The regulation of coding versus non-coding roles

A unique feature of cncRNAs that sets them apart from other RNA classes is the exquisite regulation and partitioning of their coding and non-coding functions. This can be in the form of: (1) temporal separation of the coding and non-coding activities to different developmental stages; (2) spatial segregation of the coding and non-coding roles to distinct subcellular or cellular domains; or (3) activation of one function under particular physiological/environmental conditions via specific RNA elements.

How might segregation of the coding and non-coding functions be achieved? The analysis of some cncRNAs suggests that such partitioning requires extensive transcriptional and posttranscriptional regulation of the RNAs. For instance, two major

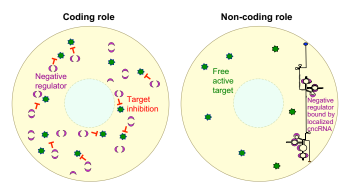


Fig. 3. cncRNAs as sequestering molecules. Structural elements in cncRNAs can sequester negative regulators. A hypothetical negative regulator (magenta half-moon) inhibits the activity of a hypothetical target (green star). However, structural features in the cncRNA (which in this case is localized to one side of the cell) allow it to bind to and sequester the negative regulator thereby enabling target activity.

SRAP isoforms of 224 or 236 amino acids, respectively, are generated by an additional upstream exon in SRA RNA that contains two initiating methionine residues, from alternative transcriptional start sites. Alternative splicing can also generate different RNA isoforms, some of which may retain the coding activity whereas others might function as non-coding RNAs. This has been observed for SRA RNA and UBE3 RNA, whereby alternative splicing leads to retention of an intron and disruption of the ORF, generating the noncoding version of these transcripts (Lanz et al., 1999; Hube et al., 2011; Valluy et al., 2015). Thus, in addition to increasing the protein-coding capacity of genomes by generating peptide isoforms, alternative splicing can also generate a variety of non-coding RNA isoforms. In addition, isoform abundance can change as development and differentiation progress, and this can influence cell fate specification. For example, the ratio of coding and noncoding SRA RNA isoforms changes during muscle cell differentiation, with myotubes expressing two to five times higher levels of non-coding SRA RNA in comparison with myoblasts. This balance between the non-coding and coding SRA RNA isoforms influences MYOD activity and myogenic differentiation (Hube et al., 2011).

Temporal segregation of the coding and non-coding activities of cncRNAs is also evident, and has been found for osk, vegt and sqt RNAs, with the non-coding activity detected at earlier developmental stages: during early oogenesis for osk, in oocytes for vegt, and in early embryos for sqt. Post-transcriptional mechanisms, such as regulated polyadenylation, splicing and translational regulation, enable this precise temporal control of RNA activity. An example of a cncRNA that undergoes extensive post-transcriptional regulation is sqt. Unprocessed sqt pre-mRNA is found in zebrafish eggs and early embryos, whereas processed sqt RNA that is poly-adenylated and spliced is detected later, from the 16-cell stage (Gore et al., 2007; Lim et al., 2012; Kumari et al., 2013). In addition, maternal sqt RNA is translationally repressed in early zebrafish embryos by the RNA-binding protein Ybx1, which interacts with the sqt 3'UTR and the eIF4E translation initiation complex. The signaling activity of Sqt/Nodal protein is detected only from the 256-cell stage (Kumari et al., 2013), suggesting that the non-coding and coding activities of sqt are temporally segregated by post-transcriptional regulation of the RNA. In Drosophila oocytes, poly-adenylation of osk RNA also stimulates Osk protein translation and is regulated by the Orb and cytoplasmic polyadenylation element binding (CPEB) proteins (Castagnetti and Ephrussi, 2003). By contrast, translation of Osk protein is repressed during early oogenesis by Bru, which binds to BREs in the osk 3' UTR sequences and recruits Cup, an eIF4E-binding protein (Nakamura et al., 2004; Chekulaeva et al., 2006). Remarkably, one cluster of BREs also mediates translational activation, and both BRE-dependent repression and activation can occur in trans, presumably by co-assembly of osk mRNAs in cytoplasmic complexes (Reveal et al., 2010).

Spatial restriction of cncRNA activity can be achieved by the localization of RNA to distinct cellular or subcellular domains, as observed for *sqt*, *osk* and *vegt*. Maternal *sqt* RNA localizes to dorsal progenitor cells by the four-cell stage in zebrafish embryos (Gore et al., 2005). Sequences in the *sqt* 3'UTR and an intact microtubule cytoskeleton are required for localization of maternal *sqt* RNA in early embryos where it carries out its non-coding role in dorsal axis formation (Gore and Sampath, 2002; Gore et al., 2005; Gilligan et al., 2011; Lim et al., 2012). Interestingly, the *sqt* dorsal localization element (DLE) overlaps with the region of the *sqt* 3' UTR that is required for Ybx1-binding, ensuring that the coding

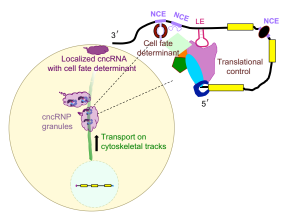


Fig. 4. cncRNAs as scaffolds. Structural elements in cncRNAs can act as scaffolds that carry ribonucleoprotein (RNP) complexes. The figure illustrates cncRNAs that act as a matrix or carrier to assemble and transport protein complexes (cncRNP granules) to specific subcellular locations via cytoskeltal transport tracks (green line). The elements that carry out the non-coding activity (NCE) are distinct from elements required for RNA localization (LE). Translational control of the cncRNA regulates the coding versus non-coding activities.

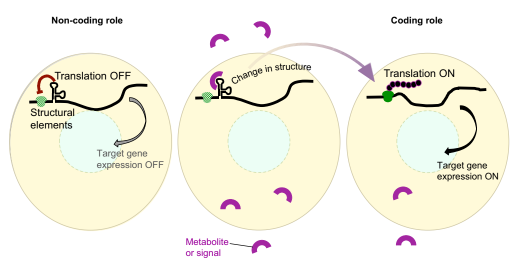
potential of localized maternal *sqt* is shut off (Gilligan et al., 2011; Kumari et al., 2013). *Drosophila osk* RNA is localized at the posterior pole of the oocyte; this localization depends on a secondary structure formed from exonic sequences in the coding region upon splicing and on Osk protein itself (Ghosh et al., 2012; Ephrussi et al., 1991; Markussen et al., 1995; Rongo et al., 1995). *vegt* RNA also shows a distinct localization pattern, being localized to the vegetal pole of *Xenopus* oocytes in a manner that depends upon sequences in the 3'UTR and the protein Igf2BP3 (also known as Vg1-RBP and Vera) (Zhang and King, 1996; Bubunenko et al., 2002; Kwon et al., 2002).

Specific regions or elements of RNA also appear to play crucial roles in determining coding versus non-coding functions. For example, the non-coding activity of *sqt* and *osk* resides in elements within the 3'UTR of these RNAs (Jenny et al., 2006; Lim et al., 2012), whereas an RNA element located between the two ORFs of *enod40* harbors its non-coding activity (Girard et al., 2003). In each of these RNAs, the coding and non-coding roles can be clearly ascribed to distinct RNA segments. However, the non-coding activity of other cncRNAs appears not to be restricted to a discrete

RNA region, but, instead, is intermingled with coding sequences and dispersed throughout the transcript, as in the case of *SRA* RNA (Lanz et al., 2002). In such RNAs, the protein-coding role may rely upon the primary sequence whereas RNA secondary/tertiary structure could engender non-coding activity. Distinguishing the coding and non-coding activities of such RNAs can be challenging.

#### **Perspectives**

The examples discussed above demonstrate that cncRNAs are emerging as key regulators of distinct developmental processes in animals and plants. It is noteworthy that all the known cncRNAs in multicellular organisms have been found in cells that are plastic, respond rapidly to their environment, and function in developmental processes (e.g. oocytes/early embryos, neurons in animals, root cells in plants). A recent study found that a significant proportion of noncoding RNAs are evolutionarily conserved and expressed in early embryos, suggesting that they might be involved in developmental processes (Necsulea et al., 2014). These observations lead to some key questions. For example, do cncRNAs represent an evolutionary link between non-coding and coding RNAs, or are they a more recent, derived group that arose independently in various clades? Which function of cncRNAs emerged first – non-coding or coding? Did these RNAs acquire new functions whilst retaining their original role? Interestingly, the non-coding RNA Xist, which mediates Xchromosome inactivation in placental mammals, is proposed to have evolved by pseudogenization of an ancestral protein-coding mRNA and loss of protein function (Duret et al., 2006), which is distinct from cncRNA genes that have either retained or acquired both coding and non-coding roles. Furthermore, it is known that riboswitches and sRNAs in bacteria act as sensors that regulate gene expression under specific environmental conditions and control key processes such as vegetative versus dormant spore formation, or motile versus sessile biofilm states (Horler and Vanderpool, 2009). This suggests that cncRNAs constitute an ancient RNA class. It is not known how many cncRNAs are present in bacterial or other genomes. Predictions based upon splice variants suggest that ~300 cncRNAs exist in the human transcriptome (Dinger et al., 2008; Ulveling et al., 2011a). However, this is likely to be an underestimate, because, in addition to intron retention and alternative splicing, other post-transcriptional mechanisms (e.g. differential poly-adenylation, RNA modifications or decorations) and differential RNA structures can also bring about cncRNA activity.



Structural elements in cncRNAs can act as sensors for metabolites, nutrients or other signals (magenta horseshoe), and regulate translation and downstream gene expression via alterations in RNA structure upon signal binding to the RNA.

Fig. 5. cncRNAs as sensors.

To identify novel cncRNAs and determine their functions, systematic analyses of mutants that specifically disrupt the transcript versus those that affect the protein-coding capacity of genes needs to be performed. Determining the features shared by dual function RNAs (similar to those found in miRNAs and long intergenic noncoding RNAs) will also facilitate the identification of novel cncRNAs. Some of these characteristics might be spatially or temporally regulated, exemplified by the presence of structural features at specific developmental stages, in certain cell types, or under particular physiological and environmental conditions. For most cncRNAs, how the features or activities of the RNAs (coding and non-coding) are regulated, and how the RNAs switch from one role to the other is largely unclear. Teasing apart the coding and noncoding activities of cncRNAs and determining how they are regulated will be challenging, but should be facilitated by mutagenesis with conventional as well as new genome editing methods.

Recent work has shown that some metabolic enzymes also have 'moonlighting' roles, functioning in their normal, wellcharacterized capacity but also in an unrelated and unexpected role. For example, in addition to its well-established function in glycolysis, the enzyme glyceraldehyde 3-phosphate dehydrogenase (GAPDH) was recently shown to have RNA-binding activity (Castello et al., 2012). In summary, these findings highlight that transcriptomes and genomes are far more complex than previously appreciated. Understanding the functions and mechanisms of action of cncRNAs will provide new insights into gene regulation. Finally, although the analysis of human diseases focuses primarily on the protein-coding capacity of the genome, it is plausible that mutations that affect non-coding functions of such cncRNAs could lead to disease states. Genome-wide analyses across phyla and throughout developmental stages, together with functional validation by classical mutagenesis, novel genome editing and RNA interrogation methods will hopefully identify novel cncRNAs and the mechanisms by which they function during plant and animal development.

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#### Competing interests

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