

# lin28 Proteins Promote Expression of 17~92 Family miRNAs During Amphibian Development

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**Background:** Lin28 proteins are post-transcriptional regulators of gene expression with multiple roles in development and the regulation of pluripotency in stem cells. Much attention has focussed on Lin28 proteins as negative regulators of *let-7* miRNA biogenesis; a function that is conserved in several animal groups and in multiple processes. However, there is increasing evidence that Lin28 proteins have additional roles, distinct from regulation of *let-7* abundance. We have previously demonstrated that lin28 proteins have functions associated with the regulation of early cell lineage specification in *Xenopus* embryos, independent of a lin28/*let-7* regulatory axis. However, the nature of lin28 targets in *Xenopus* development remains obscure. **Results:** Here, we show that mir-17~92 and mir-106~363 cluster miRNAs are down-regulated in response to lin28 knockdown, and RNAs from these clusters are co-expressed with lin28 genes during germ layer specification. Mature miRNAs derived from *pre-mir-363* are most sensitive to lin28 inhibition. We demonstrate that lin28a binds to the terminal loop of *pre-mir-363* with an affinity similar to that of *let-7*, and that this high affinity interaction requires to conserved a GGAG motif. **Conclusions:** Our data suggest a novel function for amphibian lin28 proteins as positive regulators of mir-17~92 family miRNAs. *Developmental Dynamics* 245:34–46, 2016. © 2015 The Authors. *Developmental Dynamics* published by Wiley Periodicals, Inc.

**Key words:** lin28; mir-363; let-7; mir-17~92; mir-106~363; *Xenopus*

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## Introduction

### Lin28 Proteins Are Posttranscriptional Regulators

Lin28 family proteins are posttranscriptional regulators of development and adult homeostasis. They are RNA binding proteins, characterised by a unique combination of RNA binding cold shock and zinc knuckle domains. LIN-28 was initially identified as a regulator of developmental timing in *C. elegans* and is required for the self-renewal of stem cells, with mutations in LIN-28 leading to the precocious development of late cell lineages (Moss et al., 1997; Vadla et al., 2012). Mammalian embryonic stem cells also express high levels of Lin28 proteins, which, in combination with Nanog, Oct4 and Sox2, have been used to reprogram somatic cells to a pluripotent stem cell phenotype (Viswanathan and Daley, 2010). Lin28 family genes have also been implicated as regulators in a diverse range of other biological

processes, including glucose homeostasis, tissue regeneration, and the onset of puberty in both mice and humans (Shyh-Chang and Daley, 2013; Shyh-Chang et al., 2013).

Research in several different systems has focused on the conserved role of Lin28 proteins as negative regulators of *let-7* family miRNAs. Lin28 proteins interact with both primary and precursor *let-7* miRNAs to inhibit the biogenesis of the mature biologically active forms. A prevalent model indicates an inverse relationship between levels of Lin28 proteins and mature *let-7* miRNAs (Viswanathan, 2008; Viswanathan and Daley, 2010). Typically, reduction in Lin28 function leads to increased levels of mature *let-7* miRNAs. This regulatory interaction between Lin28 proteins and *let-7* miRNAs is clearly important in multiple contexts, however, there is also increasing evidence for interactions of Lin28 proteins with a wider range of RNA targets, including other miRNA families and multiple protein coding mRNAs (Mayr and Heinemann, 2013). In the latter situation, interaction with Lin28 proteins has been shown to affect the translation of the target mRNA (Mayr and Heinemann, 2013; Shyh-Chang and Daley, 2013).

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## Lin28 Function in Amphibian Development

In an earlier study, we identified *lin28a* as a transcriptional target of FGF signalling (Branney et al., 2009). Subsequently, we investigated the function of the two Lin28-related genes, *lin28a* and *lin28b*, in *Xenopus* (Faas et al., 2013). We showed that compound knockdown of *lin28a* and *lin28b* in early development disrupts the development of axial and paraxial mesoderm. Our data indicate that *lin28* function is required in pluripotent cells of the early *Xenopus* embryo for the normal response to mesoderm inducing growth factors signals, such as FGF and activin.

## Identifying miRNA Targets of Amphibian lin28 Proteins

At present, the nature of *lin28* target RNAs in the early amphibian embryo remains elusive. Our data show that *Xenopus lin28a* and *lin28b* are able to interact with the terminal loop of *let-7* miRNAs (Faas et al., 2013). However, inhibition of *lin28* function in *Xenopus* does not lead to significant increases in the levels of mature *let-7* miRNAs in the early embryo. Therefore, the earliest perturbations in amphibian development, resulting from *lin28* knockdown, do not arise from effects on a *lin28/let-7* axis (Faas et al., 2013).

In the present study, we have undertaken a microarray based analysis to determine if other miRNAs are regulated by *lin28* in gastrula stage amphibian embryos. In contrast to the prevailing model, in which Lin28 proteins act as negative regulators of miRNA biogenesis, we find that *lin28* knockdown leads to significant down-regulation of several miRNAs. Prominent amongst these are *mir-363-5p* and *mir-363-3p*, which are derived from a common *mir-363* precursor RNA.

*mir-363* belongs to the *mir-17~92* family of miRNAs, which are encoded by the *mir-17~92*, *mir-106a~363*, and *mir-106b~25* genomic clusters. These paralogous clusters are transcribed to produce polycistronic RNAs, which are subsequently processed to form multiple, mature miRNAs with a range of related seed sequences and target specificities (Olive et al., 2010; Mogilyansky and Rigoutsos, 2013). Significantly, we find that several other miRNAs from both the *mir-17~92* and *mir-106a~363* clusters are also down-regulated in response to *lin28* inhibition, indicating that *Xenopus lin28* proteins may have a wider role in regulating the abundance *mir-17~92* family miRNAs.

We demonstrate that zygotic transcription of the *mir-17~92* and *mir-106a~363* clusters is initiated in the *Xenopus* embryo during the period of germ layer specification. We show that *mir-363-5p* and *mir-363-3p* are both expressed in the early mesoderm and later in the neuroectoderm in domains overlapping with those previously reported for *lin28a* and *lin28b* (Faas et al., 2013), suggesting a possible role for a *lin28/mir-17~92* regulatory axis in the process of germ layer specification.

The mechanism by which *lin28* proteins regulate the abundance of *mir-17~92* family miRNAs remains unclear; however, we show here that *lin28a* protein physically interacts with a GGAG motif in the terminal loop of the *pre-mir-363* miRNA. Our data support a novel function for *Xenopus lin28* proteins as positive regulators of *mir-363* miRNA abundance.

## Results

### Analysis of miRNA Abundance in *lin28* Morphant Embryos

We have previously reported the efficient knockdown of endogenous *Xenopus lin28* proteins using a combination of antisense

morpholino oligos (AMOs) directed against the three *lin28* isoforms (*lin28a1*, *lin28a2*, and *lin28b*) expressed in the embryo. In contrast to the predictions of the prevailing model for Lin28 function, we found no evidence for change in *let-7* abundance at gastrula stages following *lin28* knockdown (Faas et al., 2013). This begs the question, are there other miRNA targets of *lin28* proteins in the earliest stages of amphibian development? To begin to address this question, we have used the same AMOs to efficiently knockdown endogenous *lin28* proteins (Fig. 1A) and microarray analyses to identify changes in the abundance of miRNAs in *lin28* knockdown embryos (*lin28* morphants) at two different stages (early gastrula stage 10.5 and late gastrula stage 13). These screens were carried out using the Affymetrix microarray platform at stage 10.5 and the Exiqon microarray platform at stage 13 (Supp. Tables S1 and S2, which are available online).

An analysis of fold changes relative to controls of *Xenopus* miRNAs with an expression level of threshold  $>10$  in control and *lin28* morphant embryos at early gastrula stage 10.5 reveals several miRNAs changing in abundance. However, using a strict cut-off of  $\geq$  two-fold change and  $P \leq 0.05$ , we find that only one miRNA (*mir-363-5p*) is significantly affected in *lin28* morphants, in this case showing a 2.6-fold decrease, ( $P = 0.049$ ), indicating a positive role for *lin28* in regulating the abundance of this miRNA.

At gastrula stage 13, more miRNAs are affected in *lin28* morphants. Table 1 shows the fold changes of *Xenopus* miRNA abundance in *lin28* morphants relative to control embryos at stage 13. Only miRNAs flagged as being detected in all replicate arrays were included in this table. We find that several miRNAs show significant ( $\geq$  two-fold change and  $P \leq 0.05$ ) changes in abundance in stage 13 *lin28* morphants, including *mir-363-3p*, which, like *mir-363-5p*, is processed from a common *mir-363* precursor RNA. The *mir-363* precursor is derived from a polycistronic primary RNA which is transcribed from the *mir-106a~363* locus of clustered miRNAs; a paralogue of the well characterised *mir-17~92* miRNA cluster. Figure 1B shows the organisation of the *X. tropicalis mir-17~92* and *mir-16~363* loci as derived from the *X. tropicalis* genome sequence. As indicated in Table 1, six of eight of the significantly changing miRNAs are transcribed from these clusters, with the abundance of all decreasing in *lin28* morphants.

We next investigated quantitative changes in the levels of several *mir-17~19* and *mir-106a~363* cluster miRNAs in *lin28* morphants by quantitative real-time polymerase chain reaction (qRT-PCR). Again, we see significant decreases in the abundance of several *17~92* family miRNAs (Fig. 1C), including *mir-363-3p* and *mir-363-5p*. There is good evidence that GGAG or closely related motifs in the terminal loop regions of pre-miRNAs are important for recognition and binding by the zinc knuckle domain of *lin28* proteins (Mayr and Heinemann, 2013). Here we show that such a GGAG is present in the *Xenopus mir-363* precursor RNA. Figure 1D indicates the putative *lin28* binding motif and highlights the predicted sequences of the mature *mir-363-5p* and *mir-363-3p* miRNAs.

### Analysis of miRNA Abundance in *lin28* Over-expressing Embryos

We were interested to see how over-expression of the three *Xenopus lin28* proteins affected embryo development and the abundance of *17~92* family miRNAs. Figure 2A is a Western blot



**TABLE 1. Changes in miRNA Expression in Late Gastrula Stage 13 lin28 Morphant Embryos<sup>a</sup>**

miRNA	Mean control	Mean AMO	Fold change in morphant relative to control	Member of mir-106~363 cluster	Member of mir-17~92 cluster
<i>xtr-miR-20a</i>	1.50	0.51	-2.9		X
<i>xtr-miR-17-5p</i>	1.26	0.52	-2.4		X
<i>xtr-miR-200a</i>	1.25	0.59	-2.1		
<i>xtr-miR-20b</i>	1.29	0.61	-2.1	X	
<i>xtr-miR-301</i>	1.14	0.54	-2.1		
<i>xtr-miR-363-3p</i>	1.28	0.60	-2.1	X	
<i>xtr-miR-19a</i>	1.36	0.67	-2.0		X
<i>xtr-miR-19b</i>	1.27	0.63	-2.0	X	
<i>xtr-miR-428</i>	1.27	0.63	-2.0		
<i>xtr-miR-200b</i>	1.37	0.71	-1.9		
<i>xtr-miR-130b</i>	1.32	0.86	-1.5		
<i>xtr-miR-203</i>	1.14	0.75	-1.5		
<i>xtr-miR-30b</i>	1.13	0.90	-1.3		
<i>xtr-miR-125a</i>	1.27	1.11	-1.1		
<i>xtr-miR-126</i>	2.46	2.43	-1.0		
<i>xtr-miR-427</i>	1.09	1.04	-1.0		
<i>xtr-let-7c</i>	1.10	1.20	-0.9		
<i>xtr-let-7a</i>	1.16	1.14	1.0		
<i>xtr-let-7e</i>	0.97	0.99	1.0		
<i>xtr-miR-155</i>	1.09	1.15	1.1		
<i>xtr-miR-22</i>	0.92	1.00	1.1		
<i>xtr-miR-7</i>	0.93	1.27	1.4		

<sup>a</sup>The expression of *Xenopus* miRNAs in lin28 morphants and control embryos at late gastrula stage 13. Expression levels are shown as ratios relative to abundance in a mixed stage reference RNA sample. Only miRNAs flagged as being detected in all three replicate arrays are included. miRNAs showing  $\geq 2$  fold change and  $P \leq 0.05$  are shown in italic type. Memberships of mir-106~363 and mir-17~92 clusters are indicated.

shock and zinc-knuckle RNA binding domains, but lacks segments of N-terminal and C-terminal residues, which were predicted to be disordered (Fig. 3A). To determine if terminal truncations affect binding to canonical RNA targets, we compared the abilities of recombinant, full-length human and N- and C-terminally truncated human LIN28A (rt-LIN28A, residues 37-180) to bind the terminal loop of a *let-7-g* precursor RNA; a well characterised lin28 target, which we have previously shown to be bound by in vivo translated *Xenopus* lin28 proteins (Faas et al., 2013). Figure 3B shows that full-length and truncated human LIN28A proteins have a similar ability to bind *let-7g* RNA. In keeping with these observations, we find that Xrt-lin28a is also able to bind the *let-7g* terminal loop with high affinity ( $K_d = 314$  nM) (Fig. 3C). We provide additional evidence that Xrt-lin28a maintains its ability to discriminate genuine target RNAs. *Pre-mir-138* has previously been shown to be ineffective at competing with pre-*let-7* for binding of Lin28a protein, indicating that the terminal loop of *mir-138* is not a high affinity target of Lin28 proteins (Piskounova et al., 2008). Figure 3D shows that Xrt-lin28a protein also exhibits little binding activity toward the terminal loop of *mir-138* (L-mir-138). Even at the highest protein concentrations tested, the proportion of radiolabelled L-mir-138 bound is only 22%.

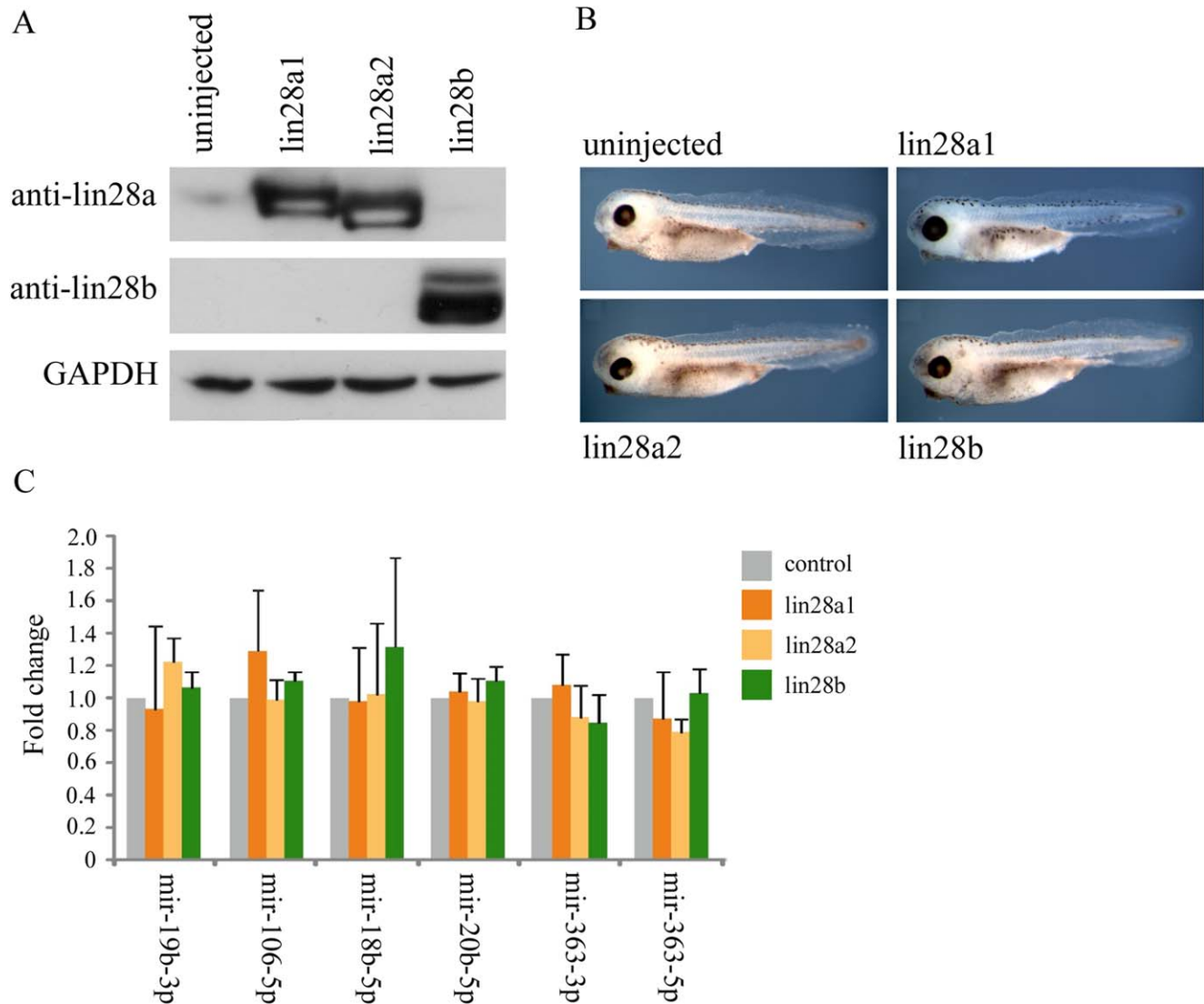
Figure 4A shows that Xrt-lin28a binds to the terminal loop region of *pre-mir-363* (L-mir-363) with a similar affinity ( $K_d = 448$  nM) to its binding with the *let-7g* terminal loop. Further evidence for the specific nature of this interaction is demonstrated by the observation that excess cold L-mir-363 competes more efficiently for binding of radiolabelled L-mir-363 to Xrt-lin28a, than does the nonrelevant L-mir-138 RNA. Thus, in the presence of a

100-fold excess of cold L-mir-363 only 3% radiolabelled L-mir-363 remains bound to Xrt-lin28a, whereas 27% remains bound in the presence of 100-fold excess of L-mir-138 RNA (Fig. 4B).

We next investigated the importance to L-mir-363 binding of the GGAG sequence motif present in the mir-363 terminal loop. A mutant *mir-363* terminal loop RNA (mL-mir-363), in which the GGAG sequence was replaced by a GUAU, was synthesised. The same substitutions have previously been shown to reduce the ability of Lin28 to bind to *let-7* (Heo et al., 2009). Figure 4C shows that Xrt-lin28a has a reduced ability to bind mL-mir-363 compared with the GGAG containing L-mir-363 RNA. Binding of radiolabelled L-mir-363 in the presence of 12.8  $\mu$ M Xrt-lin28a protein approaches 100%, whereas the binding of the mutant mL-mir-363 RNA is reduced to 58%.

### Physical Interaction of Endogenously Translated lin28a Isoforms With pre-mir-363

Our experiments using truncated recombinant *Xenopus* lin28a protein have allowed us to investigate the properties and specificity of interactions with the *mir-363* terminal loop sequence. However, it is important to note that alternative splicing of small 5' protein coding exons gives rise to two lin28a protein isoforms (lin28a1 and lin28a2) in *Xenopus* (Faas et al., 2013). To investigate binding of these two isoforms to *mir-363*, we have made use of the ability to overexpress specific proteins from injected synthetic messenger RNAs in the cells of *Xenopus* embryos. Extracts from embryos overexpressing individual isoforms can then be used as a source of full-length, in vivo translated proteins for use in EMSA



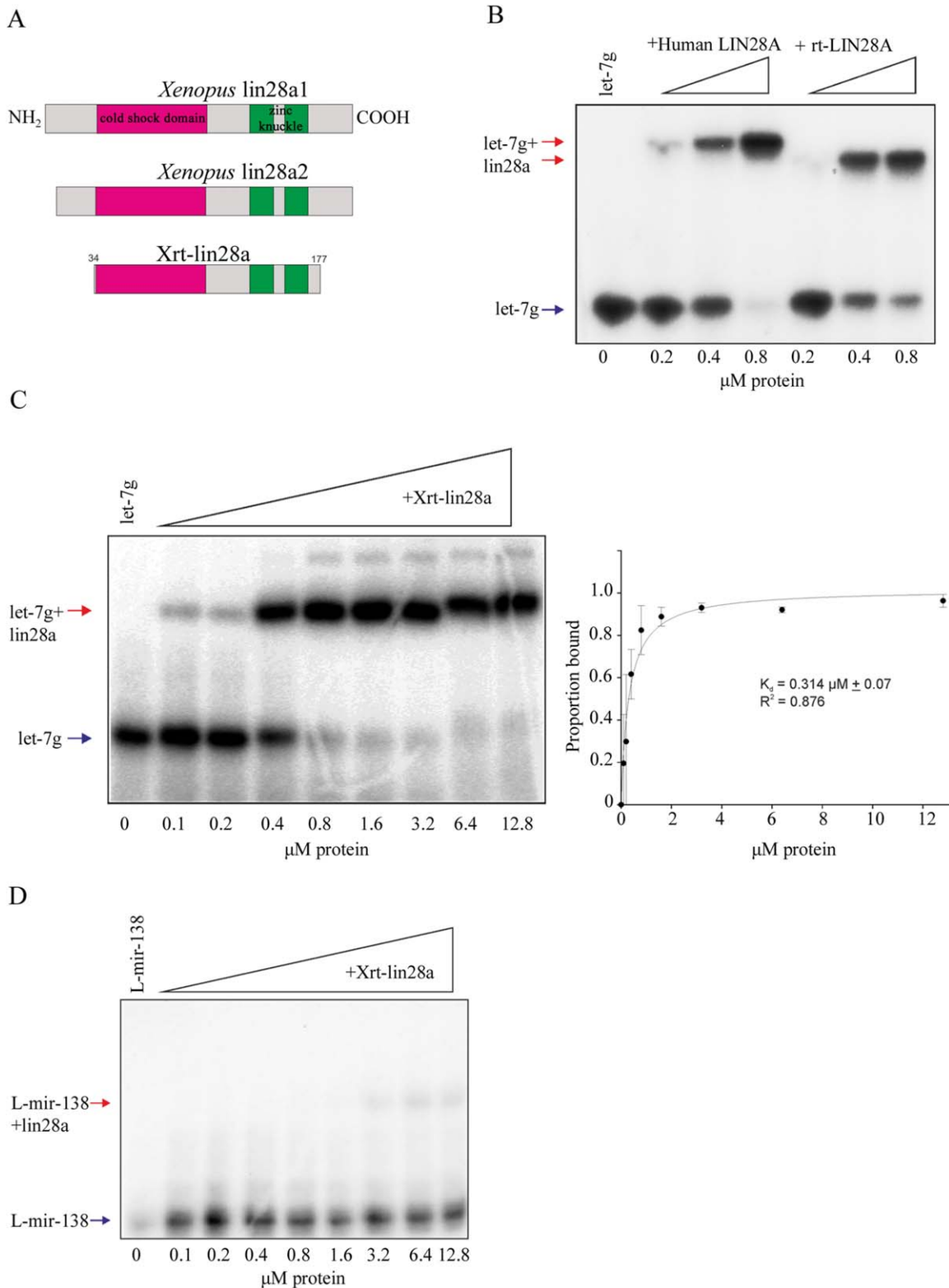
**Fig. 2.** **A:** Western blot analysis of lin28 proteins at stage 10.5 in control embryos and embryos injected with 1 ng of mRNAs coding for either lin28a1, lin28a2 or lin28b. GAPDH was used as a loading control. **B:** Phenotype of control embryos and embryos overexpressing either lin28a1, lin28a2, or lin28b at stage 38. **C:** qRT-PCR was performed on embryos injected with control embryos and either 1 ng of mRNAs coding for lin28a1, a2 or b MOs at stage 10.5. Fold change in expression of miRNAs is shown compared with controls and normalised using U6 by the  $2^{-\Delta\Delta Ct}$  method. Fold change is given as average of 3 biological replicates, with error bars representing SE.

assays. Figure 5A and 5B show that extracts from embryos overexpressing either lin28a1 or lin28a2, but not control, nonoverexpressing embryos, contain L-mir-363 binding activity. Moreover, we can attribute this binding activity to the overexpressed lin28a proteins because we are able to use an anti-lin28a antibody, but not a preimmune serum, to deplete the band corresponding to the L-mir-363+lin28 complex, giving rise to a higher molecular weight supershifted L-mir-363+lin28+Ab complex.

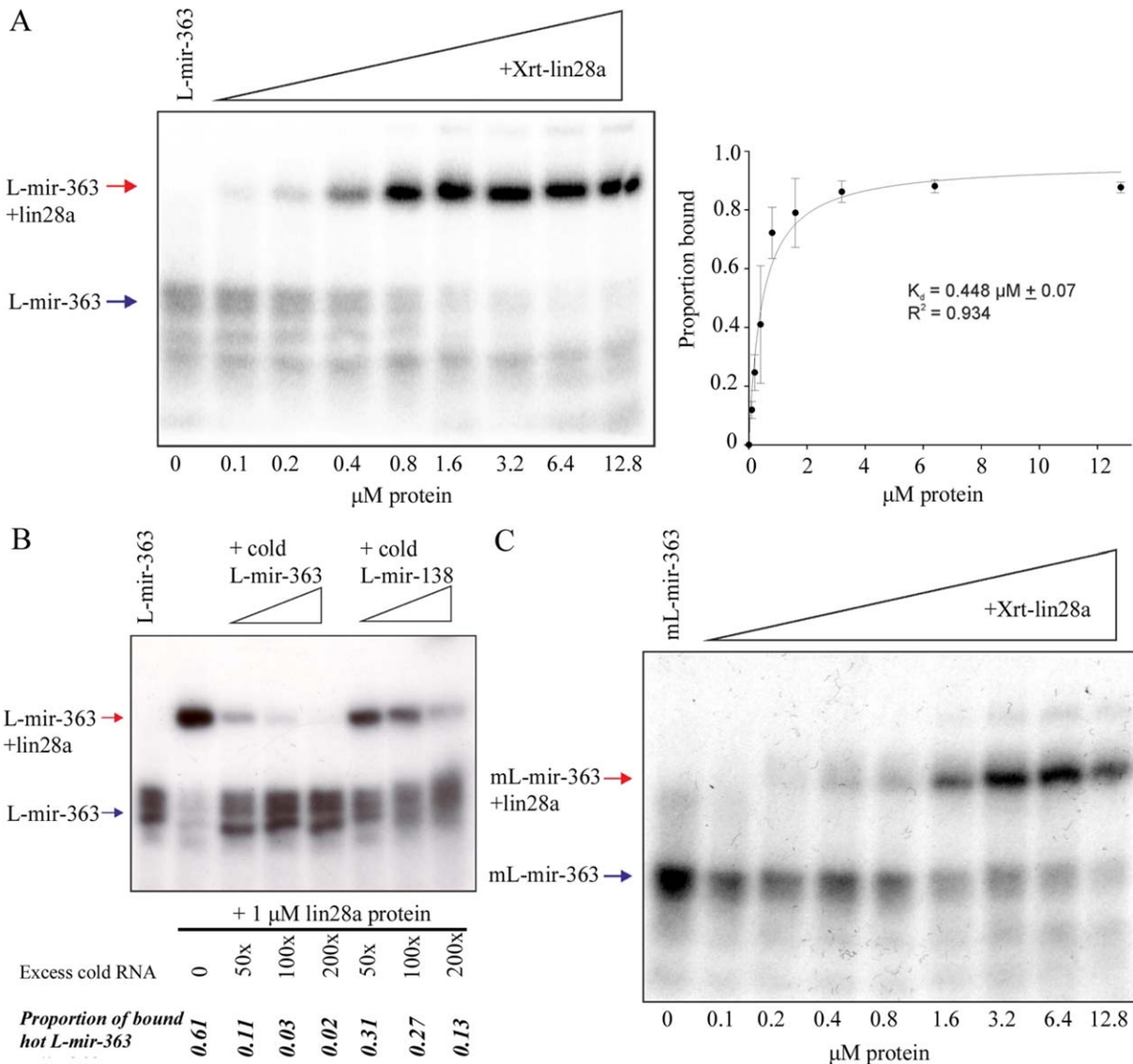
To recapitulate more accurately the binding of the full-length lin28a isoforms to the native *pre-mir-363* structure, we carried out similar binding studies with an in vitro transcribed RNA corresponding to the putative full-length *pre-mir-363*. Again we see that extracts from lin28a overexpressing embryos contain a *pre-mir-363* binding activity which can be depleted with a lin28a antibody (Fig. 5C and 5D). Interestingly, all embryo extracts contain at least two additional *pre-mir-363* binding activities (asterisks) distinct from that provided by the overexpressed lin28a proteins.

### Temporal and Spatial Expression of mir-17~92 and mir-106a~363 Clusters in the Embryo

We have provided evidence for a novel lin28 regulated pathway, involving mir-17~92 family miRNAs. However, for the proposed regulatory interactions to be relevant to normal development the components must be expressed in the same cells of the early embryo. We therefore investigated the temporal expression of the primary transcripts from the mir-17~92 cluster and mir-106a~363 clusters. Primary transcripts for both the mir-17~92 and mir-106a~363 clusters are initially detected by semi-quantitative RT-PCR at mid-blastula stage 8 (Fig. 6A). This corresponds to the time when the zygotic expression of *lin28a* is initiated (Faas et al., 2013). Figure 6B shows embryos hybridised with RNA probes designed to detect the primary transcripts from the mir-17~92 and mir-106a~363 clusters in the developing embryo. Highest levels of expression are detected in the dorsal marginal zone of the embryo at early gastrula stage 10.5.



**Fig. 3.** **A:** Scale diagram of the *Xenopus* proteins used in this study. Cold shock domains are shaded magenta and zinc knuckles green. **B:** EMSA performed with  $^{32}\text{P}$ -labelled L-let-7g and indicated concentrations of human recombinant LIN28A protein, either full-length or truncated (rt). Arrows indicate labelled RNA (blue) and LIN28A-RNA complex (red). **C:** EMSA performed with  $^{32}\text{P}$ -labelled L-let-7g and indicated concentrations of Xrt-lin28a. Gel shown is representative of  $n=3$ . Arrows indicate labelled RNA (blue) and lin28a-RNA complex (red). Band intensities were quantified from three independent experiments and the proportion bound was calculated. Data were fit by nonlinear regression as described in Materials and Methods.  $B_{\text{max}} = 1.017$ . **D:** EMSA performed with  $^{32}\text{P}$ -labelled L-mir-138 and indicated concentrations of Xrt-lin28a. Arrows indicate RNA and lin28a-RNA complex. Gel shown is representative of  $n=3$ . Arrows indicate labelled RNA (blue) and lin28a-RNA complex (red).



**Fig. 4.** **A:** EMSA performed with  $^{32}\text{P}$ -labelled L-mir-363 and indicated concentrations of Xrt-lin28a. Gel shown is representative of  $n=3$ . Band intensities were quantified from three independent experiments and the proportion bound was calculated. Data were fitted by nonlinear regression as described in *Materials and Methods*.  $B_{\text{max}} = 0.962$ . **B:** EMSA performed with  $^{32}\text{P}$ -labelled L-mir-363 and 1  $\mu\text{M}$  of Xrt-lin28a (except for RNA only lane). Arrows indicate RNA and lin28a-RNA complex. Reactions were competed with unlabelled RNA of L-mir-363 or L-mir-138 in excess levels as indicated. Band intensities were quantified and proportion of RNA bound was calculated. Gel shown is representative of  $n=2$ . Arrows indicate labelled RNA (blue) and lin28a-RNA complex (red). **C:** EMSA performed with  $^{32}\text{P}$ -labelled mL-mir-363 and indicated concentrations of Xrt-lin28a. Gel shown is representative of  $n=3$ .

### Spatial Expression of mir-363-3p and mir-363-5p in the Embryo

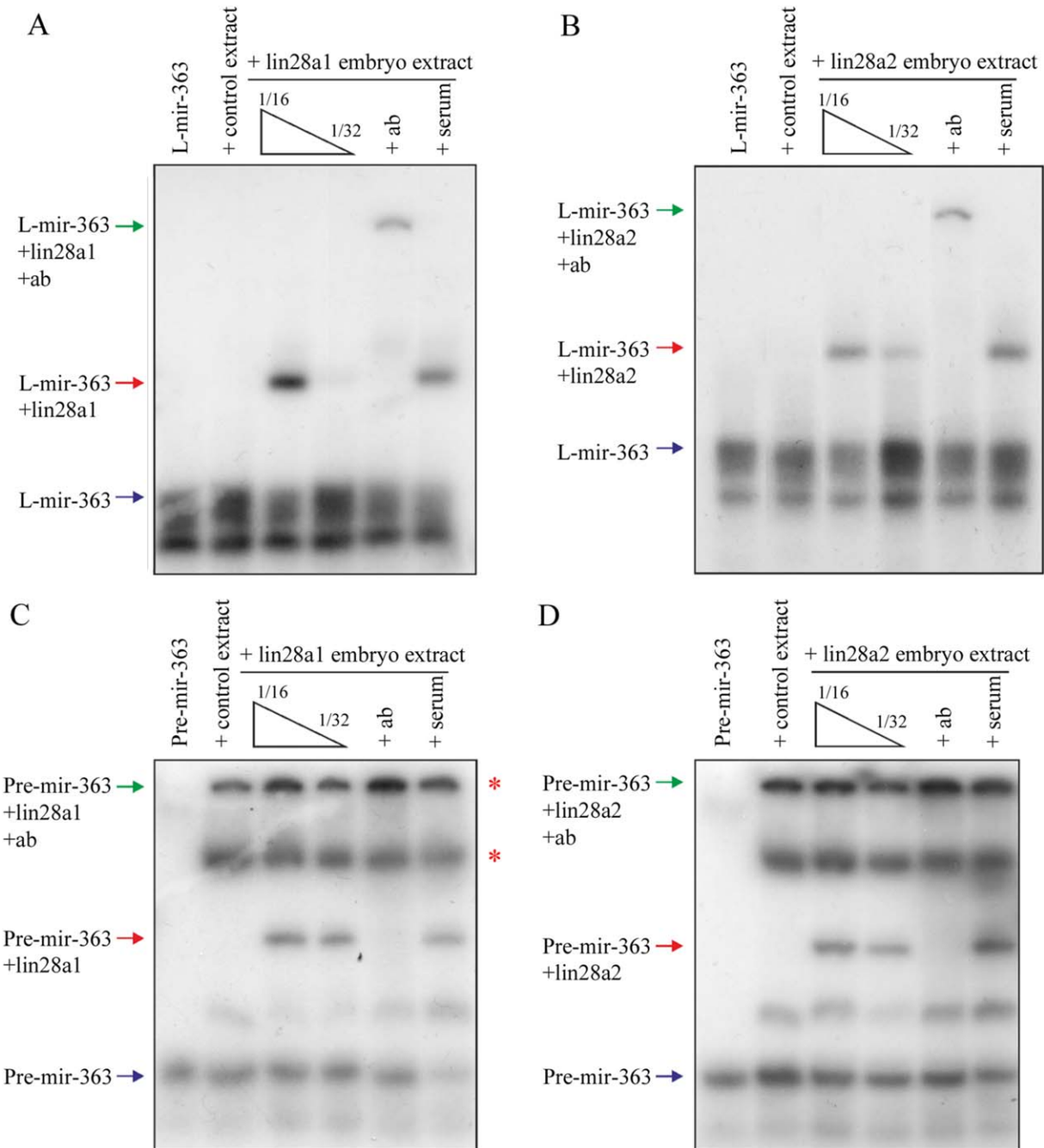
Our data provide the strongest evidence for a direct regulatory interaction between amphibian lin28 proteins and the mir-17~92 family member, *mir-363*. Anti-sense locked nucleic acid (LNA) probes contain modified nucleotides, providing increased sensitivity in detection short RNA sequences, such as miRNAs, and have previously been used to obtain highly specific in situ miRNA localisation in *Xenopus* (Sweetman et al., 2006). Figure 6C and 6D are in situ hybridisation analyses with antisense LNA probes specific for *mir-363-3p* and *mir-363-5p*. As with the primary cluster transcripts, highest levels of expression are detected in the dorsal marginal zone at the start of gastrulation. Later in devel-

opment *mir-363* miRNAs are enriched in the dorsal neural plate. Both miRNAs exhibit expression patterns similar to those previously reported for *lin28a* and *lin28b* (Faas et al., 2013).

### Discussion

#### let-7 Levels Are Unaffected in Gastrula Stage lin28 Morphants

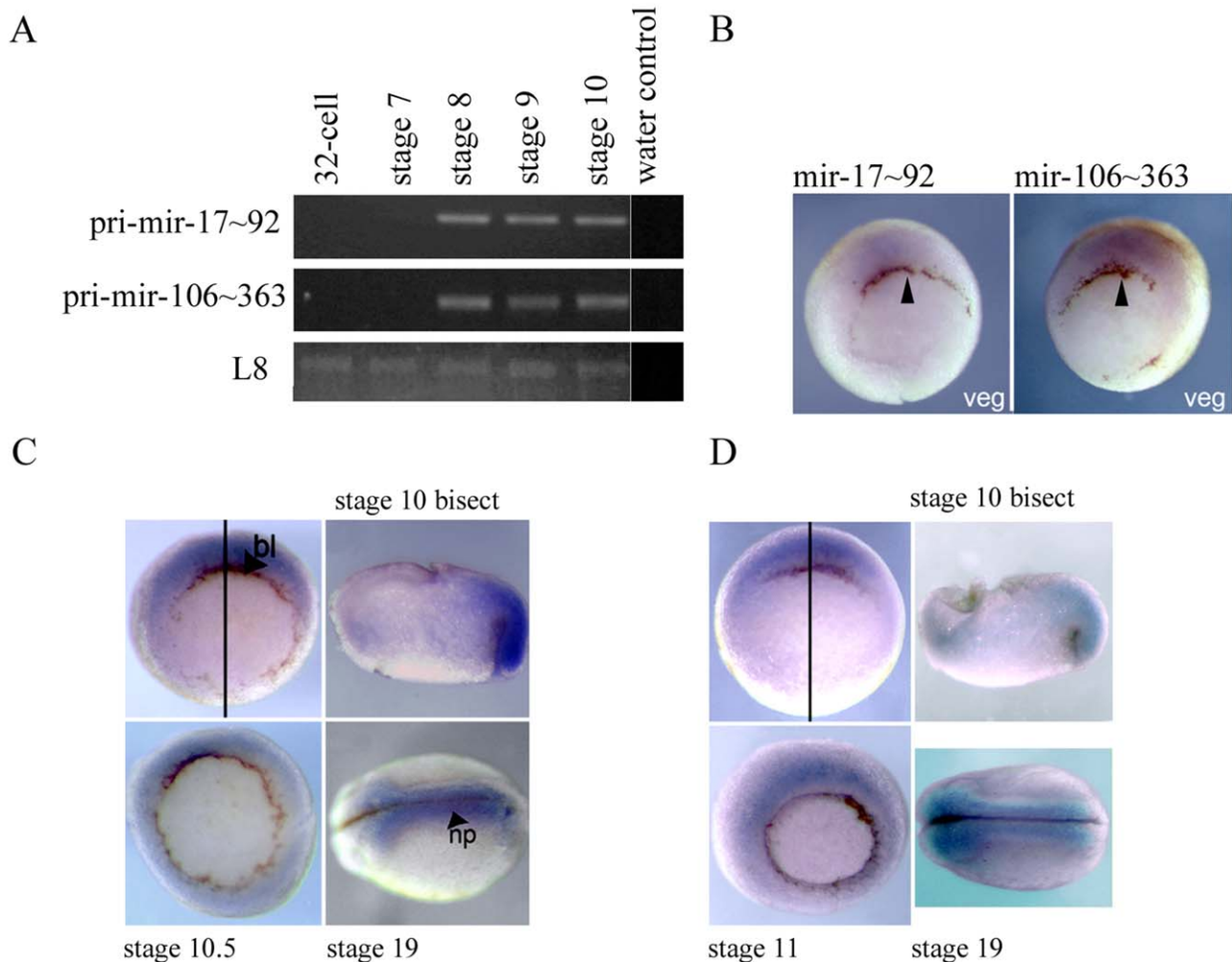
We have previously shown that lin28 function is required for the very earliest responses of pluripotent cells in the amphibian embryo to germ layer specifying growth factors. For example, levels of mesoderm lineage specific marker genes such as *brachyury*, *myoD*, and *chordin* are significantly reduced in early



**Fig. 5.** **A,B:** EMSAs performed with  $^{32}\text{P}$ -labelled L-mir-363 and embryo extract from uninjected controls or embryos injected with 1 ng of either (A) *lin28a1* or (B) *lin28a2*. Embryo extract was used at 1/16 dilution for lanes 2–3, 5–6, and at 1/32 dilution for lower concentration of overexpressing extract in lane 4. Arrows indicate unbound RNA (blue), lin28-RNA complex (red), and supershift complex of antibody-lin28-RNA (green). + Ab = 1/20 dilution  $\alpha$ -lin28a, + ser = 1/20 dilution preimmune bleed serum, both incubated with protein on ice for 20 min before addition of probe. **C,D:** EMSAs performed with  $^{32}\text{P}$ -labelled *pre-mir-363* and embryo extract from uninjected controls or embryos injected with 1 ng of either (C) *lin28a1* or (D) *lin28a2*. Embryo extract was used at 1/8 dilution for lanes 2–3, 5–6, and at 1/16 dilution for lower concentration of overexpressing extract in lane 4. Arrows indicate unbound RNA (blue), lin28-RNA complex (red), and supershift complex of antibody-lin28-RNA (green). + Ab = 1/20 dilution  $\alpha$ -lin28a, + ser = 1/20 dilution preimmune bleed serum, both incubated with protein on ice for 20 min before addition of probe.

gastrula stage compound lin28 morphants (Faas et al., 2013). Furthermore, we found that levels of mature *let-7a*, *f*, and *g* miRNAs are not significantly affected in gastrula stage lin28 morphants, leading to the proposition that, during the very earliest stages of amphibian development, lin28 proteins have functions independent of regulating *let-7* biogenesis. Here we complement and

extend this analysis using miRNA microarray-based assays, and again we detect no significant changes in the levels of any of the *let-7* family miRNAs represented on either microarray platform (*let7a*, *b*, *c*, *e*, *f*, *g*, and *i*) in gastrula stage lin28 morphants. However, we note that expression levels of *let-7* miRNAs are generally low and some family members are not detected at all (data not



**Fig. 6.** **A:** Developmental time course for expression of *pri-miR-17~92* and *pri-miR-106~363* was undertaken using RT-PCR. *L8* was used as a loading control. Image is representative of  $n=2$ . *miR-17-92* = 669 bp, *miR-106-363* = 639 bp, *L8* = 435 bp. **B:** In situ hybridisations showing expression of *pri-miR-17~92* and *pri-miR-106~363* RNAs in early development. Vegetal views of early gastrula stage 10.5 embryos, with the dorsal side is to the top. Arrows indicate the dorsal blastopore lip. **C:** In situ hybridisation using an anti-sense LNA probe showing *mir-363-3p* expression in early development. Vegetal views of gastrula stages 10 and 10.5 are shown, with dorsal-side to the top. An animal to vegetal bisect of a stage 10 embryo is shown with the animal hemisphere to the top and dorsal to the right. A dorsal view of a late neurula stage 19 embryo, anterior to the left. Plane of bisection (black line), dorsal blastopore lip (bl) and neural plate (np) are indicated. **D:** In situ hybridisation using an anti-sense LNA probe showing *mir-363-5p* expression in early development. Vegetal views of gastrula stages 10 and 11 are shown, with dorsal-side to the top. An animal to vegetal bisect of a stage 10 embryo is shown with the animal hemisphere to the top and dorsal to the right. A dorsal view of a late neurula stage 19 embryo, anterior to the left. Plane of bisection (black line) is indicated.

shown). Similar conclusions were drawn in a zebrafish *lin28* knockdown study, where no significant changes in *let-7* expression were found in morphants at 5 hpf (Ouchi et al., 2014). However, in *Xenopus* and zebrafish, increased levels of *let-7* miRNAs are detected in *lin28* morphants during later development, post-neurula stage 22 and 28 hpf, respectively, indicating that in both species there is an early *let-7*-independent, and a late *let-7*-dependent role for *lin28* proteins (Faas et al., 2013; Ouchi et al., 2014).

#### 17~92 and 106~363 Cluster miRNAs Are Down-Regulated in Gastrula Stage *lin28* Morphants

Our array analysis of morphants indicates no significant up-regulation of any miRNAs. However, several miRNAs are shown to be down-regulated, indicating a novel role for *Xen-*

*opus* *lin28* proteins as positive regulators of miRNA abundance. A notable feature of this down-regulated group is the enrichment for members of the *mir-17~92* cluster (*mir-17-5p*, *mir-19a*, and *mir-20a*) and the paralogous *mir-106~363* cluster (*mir-19b*, *mir-20b*, *mir-363-5p*, and *mir-363-3p*). These data suggest a novel regulatory interaction, where *lin28* proteins act as positive regulators of 17~92 family miRNAs.

However, we find that increasing levels of *lin28* proteins does not lead to a significant complementary up-regulation of 17~92 family miRNA abundance. This suggests that endogenous levels of *lin28* proteins are sufficient to allow maximal production of mature 17~92 family miRNAs. Thus, *lin28* levels only become limiting following knockdown. This is supported by the observations that *lin28* morphants exhibit a strong phenotype, which can be rescued by *lin28* mRNA injection (Faas et al., 2013),

whereas embryos injected with *lin28* mRNA alone develop normally (Fig. 2B).

The paralogous *mir-17~92* and *mir-106~363* clusters each code for six miRNAs, which have been highly conserved during vertebrate evolution. The *mir-17~92* cluster, in particular, has attracted a great deal of interest in recent years in relation to normal cellular function and its oncogenic potential (reviewed Mendell, 2008; Olive et al., 2010; Mogilyansky and Rigoutsos, 2013). Studies in mammals indicate that *17~92* cluster miRNA expression is high in embryonic cells, and is associated with the pluripotent state. *17~92* miRNA expression has been proposed to be part of the miRNA signature of human embryonic and induced pluripotent stem cells (Wilson et al., 2009). Mutations in the *17~92* cluster are associated with Feingold syndrome in humans, which is characterised by skeletal dysplasia (Marcelis et al., 2008). Deletion of the *17~92* locus in mice also leads to abnormal skeletal development. The phenotype of *17~92* null mice indicates additional roles in embryonic growth and morphogenesis of the heart and lungs (Ventura et al., 2008; de Pontual et al., 2011).

The *mir-106~363* cluster is less well studied. It has been reported that in mammals miRNAs from this cluster are not widely expressed and development of mice lacking the *mir-106~363* cluster is apparently normal (Ventura et al., 2008). In contrast, we find that all six *106~363* cluster miRNAs are expressed by the early amphibian embryo (Supp. Table S1). Of particular interest to the present study are *mir-363-5p* and *mir-363-3p*, which are derived from a common *mir-363* precursor RNA. Quantitative analysis of miRNA abundance in morphants indicate that, of the miRNAs analysed, *mir-363-5p* and *mir-363-3p* are most sensitive to *lin28* inhibition.

### Lin28 Proteins Bind the Terminal Loop Region of Multiple miRNAs

*Lin28* proteins physically interact with primary and precursor *let-7* miRNAs (reviewed Mayr and Heinemann, 2013). This interaction is, in part, at least, mediated by GGAG or GGAG-related sequences in the terminal loop of *pre-let-7* miRNAs, and binding provides the basis for the negative regulatory effects of *lin28* on the biogenesis of mature *let-7* miRNAs (Heo et al., 2009; Mayr and Heinemann, 2013). It is tempting to speculate that the physical interaction of *lin28* proteins with *17~92* family RNAs might also be required for the positive regulatory interaction that we report here. It has been previously reported that LIN28A is able to bind to the human *mir-363* precursor by means of a GGAG motif in its terminal loop (Heo et al., 2009). This GGAG motif is conserved in *Xenopus pre-mir-363* and both recombinant and endogenously overexpressed *lin28a* physically interact with the terminal loop sequence of *Xenopus pre-mir-363*. The affinity of this interaction is comparable with the observed for the interaction between *lin28a* and the *let-7g* terminal loop. Furthermore, we find that mutation of GGAG sequence in the *mir-363* terminal loop reduces the affinity of this interaction. In keeping with the notion that *lin28* proteins and *mir-17~92* miRNAs physically interact, we find that both *mir-363* miRNAs are expressed in similar domains to *lin28a* and *lin28b* in the presumptive mesoderm of gastrula stage *Xenopus* embryos (Faas et al., 2013).

*Lin28* proteins inhibit the biogenesis of *let-7* miRNAs using multiple mechanisms. One reported mechanism requires LIN28A dependent recruitment of the Tut4 enzyme to a *pre-let-7* containing complex and subsequent Tut4 mediated polyuridylation and

inactivation of *pre-let7* miRNAs (Heo et al., 2008, 2009). Several other human miRNAs have been shown to contain a terminal loop GGAG motif and are bound by LIN28A; however, this association does not always lead to Tut4 mediated polyuridylation and destabilisation (Heo et al., 2009). Thus, *Lin28* binding to a GGAG motif in the terminal loop can have different consequences, depending on the target miRNAs involved.

At present, we do not know the mechanism by which amphibian *lin28* proteins promote the expression of *17~92* family miRNAs in the early embryo. Indeed, our data do not rule out the possibility of *lin28* proteins, indirectly or directly, regulating *17~92* miRNA expression by multiple mechanisms. However, an attractive hypothesis is that the binding of *lin28* to the terminal loop region of the *pre-mir-363* sequence within the *106~363* polycistron somehow promotes subsequent processing to the precursor and mature miRNAs derived from the primary transcript. In regard to the related *17~92* polycistron, we have not yet investigated physical interactions with *lin28* proteins. We have not identified GGAG-like sequences in the terminal loop region of *17~92* polycistron derived pre-miRNAs; however, there are multiple GGAG motifs present within the intergenic regions of the polycistronic primary transcript (data not shown). It will be interesting to determine whether these motifs can act as binding sites for *lin28* proteins.

### Lin28, *let-7* Family, and *17~92* Family; Key Components of a Pluripotency Network

*Lin28* expression is associated with pluripotent mammalian stem cells in culture and pluripotent cells in the early amphibian embryo that respond to the earliest lineage specifying growth factor signals. A key function of *Lin28* proteins in mammalian stem cells is to inhibit the biogenesis of *let-7* miRNAs. During differentiation of stem cells, *lin28* levels fall and levels of biologically active *let-7* miRNAs rise (Viswanathan et al., 2008; Viswanathan and Daley, 2010). While a role for *lin28* regulating *let-7* biogenesis in postneurula stage amphibian embryos is supported, there is no evidence that this function is important in very early development, perhaps because transcription of *let-7* miRNAs is low. In contrast to *let-7* miRNAs, elevated expression of *17~92* family miRNAs is associated with the pluripotent state. It is tempting to speculate that in some pluripotent cell populations *lin28* proteins might play a dual role in inhibiting *let-7* and promoting *17~92* family expression. We note that similar dual, opposite effects on the regulation of *let-7* and *17~92* miRNAs have also been reported for the hnRNPA1 RNA binding protein, which like *Lin28* inhibits *let-7* biogenesis, but promotes the biogenesis of the *17~92* cluster miRNA, *mir-18a* (Guil and Caceres, 2007; Michlewski and Caceres, 2010).

Taken together, our results suggest a novel regulatory function for *lin28* proteins in the pluripotent cells of the early amphibian embryo, in which *lin28* proteins positively regulate levels of mature *17~92/106~363* cluster miRNAs in the early embryo, which contrasts with their activity of negatively regulating mature *let-7* miRNA levels in mammalian stem cell populations.

## Experimental Procedures

### Embryo Methods

*Xenopus tropicalis* embryos were produced as previously described (Khokha et al., 2005; Winterbottom et al., 2010).

Embryos were injected at two- or four-cell stage and cultured at 22 °C.

Samples for miRNA analysis were isolated using the miRVana miRNA isolation kit (Applied Biosystems). The protocol was carried out according to manufacturer's instructions with the modification that after lysis samples were centrifuged for 10 min at 4 degC and supernatant removed to a fresh tube.

### Western Bot Analysis

Western blots were carried out as previously described, using affinity-purified *X. tropicalis* anti-lin28 antisera raised by inoculation of peptides corresponding to the C-terminal sequences of *X. tropicalis* lin28a1/a2 (EEQPISEEQELIPETME) or lin28b (SRKGPSVQKRKKT) proteins (Faas et al., 2013).

### Knockdown of lin28a and lin28b

Compound knockdown of lin28a1+a2+b was accomplished using a total of 25 ng per embryo of a mixture containing 10 ng lin28a1 + 10 ng lin28a2 + 5 ng lin28b AMOs (Gene Tools, LLC), as previously described (Faas et al., 2013). Injections were carried out into all cells at either the two- or four-cell stage, with a maximum of 10 nl/embryo. Injections were targeted to the marginal zone.

### Overexpression of lin28a and lin28b

The coding regions of *lin28a1*, *lin28a2*, and *lin28b* were PCR amplified and sub-cloned into the Cs2 + mRNA transcription vector. mRNA synthesis was as previously described (Branney et al., 2009). All cells were injected at the two- or four-cell stage, with a total of 1 ng/embryo of each mRNA. Injections were targeted to the marginal zone.

### Affymetrix miRNA Array Analysis

RNA was isolated from control embryos and knockdown as described above at early gastrula stage 10.5. The quality of the RNA was verified using the Agilent 2011 Bioanalyzer (Agilent). Samples were processed in the University of York, Department of Biology Technology Facility. One microgram samples were processed in the University of York, Department of Biology Technology Facility. RNA was labelled using HSR FlashTag Biotin RNA labelling kit (Genisphere) according to manufacturer's instructions, which included the addition of spike-in RNA controls to act as a method control. Samples were then hybridised to Genechips miRNA 2.0 (Affymetrix) overnight, and washed on a Fluidics Station 450 (Affymetrix), all carried out according to manufacturer's instructions. Scanning of the chips was carried out using an Affymetrix Genechip Scanner. CEL files were processed using Affymetrix QC tools software to provide background detection and quantile normalisation with a final median polish and log transformation. *Xenopus* feature data were extracted and statistical comparisons undertaken using a paired, two-tail Student's *t*-test. The complete triplicate summarization data set for the *Xenopus* features are shown in Supplementary Table S1. These data have been deposited in the ArrayExpress Archive (<https://www.ebi.ac.uk/arrayexpress/>) with accession number E-MTAB-3936.

### Exiqon miRNA Array Analysis

Compound knockdown of lin28a1+a2+b was carried out as described above. Control embryos were injected with 30 ng of a standard control MO. RNA was isolated from experimental embryos at late gastrula stage 13. Quality control, sample processing, and preliminary data processing, including normalization were undertaken as a service by Exiqon A.S. Expression levels were calculated as fold changes relative to a mixed stage reference RNA sample. *Xenopus* feature data were extracted and statistical comparisons were undertaken using a paired, 2-tail Student's *t*-test. The median log ratios for the triplicate *Xenopus* data set are shown in Supplementary Table S2. These data have been deposited in the ArrayExpress Archive (<https://www.ebi.ac.uk/arrayexpress/>) with accession number E-MTAB-3939.

### Semi-quantitative PCR Analysis of miRNA Cluster Primary Transcript Abundance

Total RNA was extracted using TRI reagent (Sigma) according to the manufacturer's instructions. An additional precipitation step was undertaken using 7.5 M LiCl and 0.05 M EDTA at -80 degC overnight. cDNA was synthesised from total RNA using 1 µg RNA random hexamers (Invitrogen) and SuperScript II Reverse Transcriptase (Invitrogen) according to manufacturer's instructions. cDNA was diluted 1/5 for use in RT-PCR reactions using PCR Master Mix (Promega). Primer sequences are shown below.

L8 forward:	GGGCTRTC GACTTYGCTGAA
L8 reverse:	ATACGACCACCWCCAGCAAC
miR-17~92 cluster forward:	TGCAGTGAAGGCACTTGTAG
miR-17~92 cluster reverse:	TAAACAGGCCGGGACAAG
mir-106a~363 cluster forward:	TGCTGGACACCTGTACT
mir-106a~363 cluster reverse:	TTCTGCGTTTACAGATGGA

### miRNA qRT-PCR

Samples to be used for miRNA analysis were isolated using the miRVana miRNA isolation kit (Applied Biosystems) as described above. cDNA was synthesised from 10 ng RNA/RT reaction with miRNA-specific primers for TaqMan assays (Applied Biosystems) using the TaqMan MicroRNA Reverse Transcription Kit (Applied Biosystems) as manufacturer's instructions. qRT-PCR was carried out using TaqMan Universal Master Mix II (Applied Biosystems) with Taqman miRNA probes (Applied Biosystems) according to manufacturer's instructions. All reactions were performed in quadruplicate per sample on an ABI Prism 7000 detection system (Applied Biosystems) with thermal cycling at 95 degC for 10 min, followed by 40 cycles of 95 degC for 15 sec and 60 degC for 1 min. Gene expression levels were normalised to U6 snRNA using the  $2^{-\Delta\Delta Ct}$  method. Preliminary experiments had shown that U6 snRNA was a suitable control for this purpose (data not shown). Assays used were: hsa-miR-19b, hsa-miR-20b, hsa-miR-363#, hsa-miR-363, hsa-miR-18b, hsa-let-7a, hsa-let-7f, custom xtr-mir-106a (Applied Biosystems). It is important to note that the inclusion of stem loop structures in the primers used in the miRNA specific cDNA syntheses allow for the detection of mature, biologically active miRNAs.

## Whole-Mount In Situ Hybridisation for miRNA Cluster Primary Transcripts

To generate whole-mount in situ hybridisation probes for the miR-17~92 and mir-106a~363 clusters, cDNAs corresponding to sections of the miR-17~92 and mir-106a~363 primary transcripts were cloned in the pGEM®-T Easy vector following PCR amplification using *X. tropicalis* genomic DNA as template and the following primers.

mir-17~92 cluster forward:	TGCAGTGAAGGCACTTGTAG
mir-17~92 cluster reverse:	TAAACAGGCCGGACAAG
mir-106a~363 cluster forward:	TGCTGGACACCTGTACT
mir-106a~363 cluster forward:	TTCTGCGGTTTACAGATGGA

Digoxigenin (DIG) -labelled antisense in situ probes were transcribed and in situ hybridisation was carried out as previously described (Harland, 1991) with slight modification (Reece-Hoyes et al., 2002).

## Whole-Mount In Situ Hybridisation for miRNA

Probes used were 5'-DIG labelled LNA miRNA detection probes (Exiqon), named "hsa-miR-363-3p," and "xtr-miR-363\*." Protocol was carried out as described previously (Sweetman, 2011), with modifications advised by Grant Wheeler (University of East Anglia, UK, personal communication). Probes were preabsorbed six times by hybridising with the probe overnight against stage 35 embryos. Colour development was with NBT/BCIP substrate. When signal began to develop, embryos were washed at 4 degC overnight and subjected to repeat cycles of colour reaction and washes until a strong specific signal. Embryos were then fixed and bleached with hydrogen peroxide to remove pigment before photography.

## Recombinant Lin28 Protein Production

Relevant coding sequences were cloned into the pET28a expression vector. Recombinant Lin28 proteins were expressed overnight at 16 degC in B834 *E. coli* cells grown in LB media, following induction with 1 mM IPTG. Cells were harvested by centrifugation and pellets resuspended in a solution containing either 50 mM sodium phosphate pH 7.8, 250 mM NaCl 1 mM DTT, 20 mM imidazole, and 10% w/v glycerol (rt-LIN28A); or 50 mM Tris HCl pH 7.5, 500 mM NaCl, 0.5 mM DTT, 20 mM imidazole, and 10% w/v sucrose (Xrt-lin28a). The resuspension solution was supplemented with 0.5 µg/ml leupeptin, 0.7 µg/mL pepstatin, and 1 mM AEBSF protease inhibitors. Cells were lysed by sonication and the lysate applied to a 5 ml HisTrap column (GE Healthcare). Bound protein was eluted using a linear imidazole gradient (20–500 mM). Fractions containing Lin28 were analysed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE), pooled, concentrated, and applied to an S200 gel filtration column (GE Healthcare) in a running buffer consisting of either 20 mM Tris pH 7.5, 150 mM NaCl, 1 mM DTT, 10% w/v glycerol (rt-LIN28A), or 10 mM Tris pH 7.5, 150 mM NaCl, 2 mM DTT, and 10% w/v sucrose (Xrt-lin28a). Eluting fractions containing Lin28 were analyzed by SDS-PAGE, pooled and concentrated, before being flash frozen in liquid N<sub>2</sub> and stored at -80 degC. Proteins

were produced as N-terminal fusions with the sequence, MGSSHHHHHSSGLVPRGSHM, containing a His-tag and thrombin digest site. In the case of the human rt-Lin28a, the N-terminal His-tag was removed by thrombin digest before gel filtration.

## Full-Length Human LIN28A Protein

MGSVSNQQFAGGCAKAAEEAPEEAPEDAARAADPEQLLHGAGICKWFNVRMFGFLSMTARAGVALDPPVDVVFVHQSKLHMEGFRSLKEGEAVEFTFKKSAKGLSIRVTGPGGVFCIGSERRPKGKSMQKRRSKGDRCYNCGLDHHAKECKLPPQPKKCHFCQSISHMVASCPLKAQQGPSAQGKPTYFREEEEEIHSPTLLPEAQN

## N- and C-Ternally Truncated Human LIN28A Protein (rt-LIN28A), Residues 37-180

LLHGAGICKWFNVRMFGFLSMTARAGVALDPPVDVVFVHQSKLHMEGFRSLKEGEAVEFTFKKSAKGLSIRVTGPGGVFCIGSERRPKGKSMQKRRSKGDRCYNCGLDHHAKECKLPPQPKKCHFCQSISHMVASCPLKAQQ

## N- and C-Ternally Truncated *Xenopus* lin28a Protein (Xrt-lin28a), Residues 34-177

MGSSHHHHHSSGLVPRGSHMGSVGVCKWFNVRMFGFLTMTKKEGTDLETPVDVVFVHQSKLHMEGFRSLKEGESVEFTFKKSSKGLSIRVTGPGGAPCIGSERRPKVKGQKRRQKGDRCYNCGLDHHAKECKLPPQPKKCHFCQSPNHMVAQCPAKASQAAN. (Leader containing the His-tag and thrombin cleavage site is indicated in bold.)

## RNA EMSAs

Pre-cursor *mir-363* RNA was synthesised in vitro. DNA templates for *pre-mir-363* were produced by PCR, to include an SP6 RNA polymerase promoter at the beginning of the sequence, using a plasmid containing the *Xenopus* mir-106a-363 as template and the following primers. RNA was synthesised using SP6 Megascript kit (Ambion) according to manufacturer's instructions.

### *Pre-mir-363*

Forward:	Reverse:
ATTTAGGTGACAC	TAGGCAAGGCAGTGG
TATAGGGCTGAGG	CCTGTACAG
TAGTTGTTT	

RNA oligonucleotides used in RNA mobility shift assays were synthesised by Dharmacon.

### L-mir-138

UUGUGAAUCAGGCCGUGACCACUCAGAAAACGGCUACUUCA CAAC

### L-mir-363

UGCAAUUUUUUUUAGUUUGGUAGGAGAAAAUUUGCmL-mir-363

UGCAAUUUUUUUUAGUUUGGUAUGAUAAAAUUUGC

RNA oligonucleotides and mir-RNA precursors were radioactively labelled with <sup>32</sup>P ATP using the KinaseMax kit (Ambion) according to manufacturer's instructions.

Recombinant protein EMSAs were performed with the proteins described above. For embryo extract EMSAs, uninjected *X. laevis* controls embryos and embryos injected with 1 ng *lin28a1*,

*lin28a2*, or *lin28b* mRNA (Faas et al., 2013) were lysed in 50 mM Tris-HCl pH 7.9, 25% glycerol, 50 mM KCl, 2 mM DTT, 0.1 mM EDTA, 1/100 Protease inhibitor cocktail III [Calbiochem] at 10  $\mu$ l/embryo. Lysates were cleared by centrifugation and extracts were diluted as required in the lysis buffer.

Binding reactions were carried out as described previously (Piskounova et al., 2008). Labelled RNA probes were incubated with protein in binding buffer (60 mM KCl, 10 mM HEPES, pH 7.6, 3 mM MgCl<sub>2</sub>, 5% glycerol, 1 mM DTT, 5  $\mu$ g/ $\mu$ l heparin [Sigma], and 150 ng yeast total RNA competitor [Ambion]) for 30 min at room temperature.

The custom anti-*lin28a* antibody (Enzo Life Sciences (UK) Ltd), used for the embryo extract supershift assays, has been previously described (Faas et al., 2013) and was used at 1/20 dilution per binding reaction, with 1/20 dilution preimmune bleed used as a serum control. 20 units of RNasin (Promega) were added per binding. Antibody was preincubated with protein and binding buffer for 20 min on ice, before labelled probe was added for a further 20 min at room temperature.

Samples were run on a 10% native polyacrylamide gel. Gels were dried and exposed either to a Phosphor Screen (GE Healthcare) and were scanned, processed and analysed using a Bio-Rad Molecular FX Imager and Quantity One software (Bio-Rad); or exposed to Hyperfilm ECL film (Amersham) and films analysed using Image J. The proportion of RNA bound at each protein concentration was calculated, and the  $K_d$  determined by nonlinear regression using the SigmaPlot software package, with the equation:

$$\text{Proportion bound} = \frac{B_{\max} [\text{lin28}]}{K_d + [\text{lin28}]}$$

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## References

- Branney PA, Faas L, Steane SE, Pownall ME, Isaacs HV. 2009. Characterisation of the fibroblast growth factor dependent transcriptome in early development. *PLoS One* 4:e4951–e4951.
- de Pontual L, Yao E, Callier P, Faivre L, Drouin V, Cariou S, Van Haeringen A, Genevieve D, Goldenberg A, Oufadem M, Manouvrier S, Munnich A, Vidigal JA, Vekemans M, Lyonnet S, Henrion-Caude A, Ventura A, Amiel J. 2011. Germline deletion of the miR-17 approximately 92 cluster causes skeletal and growth defects in humans. *Nat Genet* 43:1026–1030.
- Faas L, Warrander FC, Maguire R, Ramsbottom SA, Quinn D, Genevieve P, Isaacs HV. 2013. Lin28 proteins are required for germ layer specification in *Xenopus*. *Development* 140:976–986.
- Guil S, Caceres JF. 2007. The multifunctional RNA-binding protein hnRNP A1 is required for processing of miR-18a. *Nat Struct Mol Biol* 14:591–596.
- Harland RM. 1991. In situ hybridization: an improved whole-mount method for *Xenopus* embryos. *Methods Cell Biol* 36:685–695.
- Heo I, Joo C, Cho J, Ha M, Han J, Kim VN. 2008. Lin28 mediates the terminal uridylation of let-7 precursor MicroRNA. *Mol Cell* 32:276–284.
- Heo I, Joo C, Kim YK, Ha M, Yoon MJ, Cho J, Yeom KH, Han J, Kim VN. 2009. TUT4 in concert with Lin28 suppresses microRNA biogenesis through pre-microRNA uridylation. *Cell* 138:696–708.
- Khokha MK, Yeh J, Grammer TC, Harland RM. 2005. Depletion of three BMP antagonists from Spemann's organizer leads to a catastrophic loss of dorsal structures. *Dev Cell* 8:401–411.
- Marcelis CL, Hol FA, Graham GE, Rieu PN, Kellermayer R, Meijer RP, Lugtenberg D, Scheffer H, van Bokhoven H, Brunner HG, de Brouwer AP. 2008. Genotype-phenotype correlations in MYCN-related Feingold syndrome. *Hum Mutat* 29:1125–1132.
- Mayr F, Heinemann U. 2013. Mechanisms of Lin28-mediated miRNA and mRNA regulation—a structural and functional perspective. *Int J Mol Sci* 14:16532–16553.
- Mendell JT. 2008. miRiad roles for the miR-17-92 cluster in development and disease. *Cell* 133:217–222.
- Michlewski G, Caceres JF. 2010. Antagonistic role of hnRNP A1 and KSRP in the regulation of let-7a biogenesis. *Nat Struct Mol Biol* 17:1011–1018.
- Mogilyansky E, Rigoutsos I. 2013. The miR-17/92 cluster: a comprehensive update on its genomics, genetics, functions and increasingly important and numerous roles in health and disease. *Cell Death Differ* 20:1603–1614.
- Moss EG, Lee RC, Ambros V. 1997. The cold shock domain protein LIN-28 controls developmental timing in *C. elegans* and is regulated by the *lin-4* RNA. *Cell* 88:637–646.
- Olive V, Jiang I, He L. 2010. mir-17-92, a cluster of miRNAs in the midst of the cancer network. *Int J Biochem Cell Biol* 42:1348–1354.
- Ouchi Y, Yamamoto J, Iwamoto T. 2014. The heterochronic genes *lin-28a* and *lin-28b* play an essential and evolutionarily conserved role in early zebrafish development. *PLoS One* 9:e88086.
- Piskounova E, Viswanathan SR, Janas M, LaPierre RJ, Daley GQ, Sliz P, Gregory RI. 2008. Determinants of microRNA processing inhibition by the developmentally regulated RNA-binding protein Lin28. *J Biol Chem* 283:21310–21314.
- Reece-Hoyes JS, Keenan ID, Isaacs HV. 2002. Cloning and expression of the *Cdx* family from the frog *Xenopus tropicalis*. *Dev Dyn* 223:134–140.
- Shyh-Chang N, Daley GQ. 2013. Lin28: primal regulator of growth and metabolism in stem cells. *Cell Stem Cell* 12:395–406.
- Shyh-Chang N, Zhu H, Yvanka de Soysa T, Shinoda G, Seligson MT, Tsanov KM, Nguyen L, Asara JM, Cantley LC, Daley GQ. 2013. Lin28 enhances tissue repair by reprogramming cellular metabolism. *Cell* 155:778–792.
- Sweetman D. 2011. In situ detection of microRNAs in animals. *Methods Mol Biol* 732:1–8.
- Sweetman D, Rathjen T, Jefferson M, Wheeler G, Smith TG, Wheeler GN, Munsterberg A, Dalmay T. 2006. FGF-4 signaling is involved in mir-206 expression in developing somites of chicken embryos. *Dev Dyn* 235:2185–2191.
- Vadla B, Kemper K, Alaimo J, Heine C, Moss EG. 2012. *lin-28* controls the succession of cell fate choices via two distinct activities. *PLoS Genet* 8:e1002588.
- Ventura A, Young AG, Winslow MM, Lintault L, Meissner A, Erkeland SJ, Newman J, Bronson RT, Crowley D, Stone JR, Jaenisch R, Sharp PA, Jacks T. 2008. Targeted deletion reveals essential and overlapping functions of the miR-17 through 92 family of miRNA clusters. *Cell* 132:875–886.
- Viswanathan SR, Daley GQ, Gregory RI. 2008. Selective blockade of microRNA processing by Lin28. *Science (New York, N.Y.)* 320:97–100.
- Viswanathan SR, Daley GQ. 2010. Lin28: a microRNA regulator with a macro role. *cell* 140:445–449.
- Viswanathan SR, Daley GQ, Gregory RI. 2008. Selective blockade of microRNA processing by Lin28. *Science* 320:97–100.
- Wilson KD, Venkatasubrahmanyam S, Jia F, Sun N, Butte AJ, Wu JC. 2009. MicroRNA profiling of human-induced pluripotent stem cells. *Stem Cells Dev* 18:749–758.
- Winterbottom EF, Illes JC, Faas L, Isaacs HV. 2010. Conserved and novel roles for the Gsh2 transcription factor in primary neurogenesis. *Development* 137:2623–2631.