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Independent regulation of vertebral number and vertebral identity by microRNA-196 paralogs

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The *Hox* genes play a central role in patterning the embryonic anterior-to-posterior axis. An important function of *Hox* activity in vertebrates is the specification of different vertebral morphologies, with an additional role in axis elongation emerging. The miR-196 family of microRNAs (miRNAs) are predicted to extensively target *Hox* 3' UTRs, although the full extent to which miR-196 regulates *Hox* expression dynamics and influences mammalian development remains to be elucidated. Here we used an extensive allelic series of mouse knockouts to show that the miR-196 family of miRNAs is essential both for properly patterning vertebral identity at different axial levels and for modulating the total number of vertebrae. All three miR-196 paralogs, 196a1, 196a2, and 196b, act redundantly to pattern the midthoracic region, whereas 196a2 and 196b have an additive role in controlling the number of rib-bearing vertebra and positioning of the sacrum. Independent of this, 196a1, 196a2, and 196b act redundantly to constrain total vertebral number. Loss of miR-196 leads to a collective up-regulation of numerous trunk *Hox* target genes with a concomitant delay in activation of caudal *Hox* genes, which are proposed to signal the end of axis extension. Additionally, we identified altered molecular signatures associated with the *Wnt*, *Fgf*, and *Notch/segmentation* pathways and demonstrate that miR-196 has the potential to regulate *Wnt* activity by multiple mechanisms. By feeding into, and thereby integrating, multiple genetic networks controlling vertebral number and identity, miR-196 is a critical player defining axial formulae.

Hox genes | microRNA | miR-196 | axial patterning | vertebral specification

A defining feature of the vertebrate animals is the presence of a segmented vertebral column. Species are uniquely characterized by the total number of vertebrae that form and by the regionalization of these vertebra along the anterior-to-posterior axis into groups with distinct morphologies (e.g., cervical, thoracic, lumbar, and sacral). The genetic determinants of vertebral number and vertebral identity have largely been considered as separate; thus how, or even whether, these processes are molecularly integrated remains to be clearly elucidated.

Vertebral precursors, known as somites, arise by continued expansion and segmentation of a region of the caudal embryo, the presomitic mesoderm (PSM) (1). Expansion of the PSM requires a self-renewing axial progenitor population that initially resides in the node-streak border of the epiblast and subsequently repositions to the tailbud (2–6). These progenitors provide a source of cells that, following ingress through the primitive streak, populate the PSM and other derivatives to drive posterior elongation. Key players in this process include genes involved in *Wnt* and *Fgf* signaling, in addition to the *Cdx* transcription factors, as evidenced by severe axis truncations when each are mutated (7). Balancing the expansion of this cell population, cells of the anterior PSM bud off to form somites with a rhythmic periodicity inherent to each species. The eventual exhaustion of progenitor self-renewal capacity is thought to halt axis elongation, the timing

of which is a critical factor in establishing species-specific vertebral number (8).

Within vertebral precursors, specific combinations of *Hox* transcription factors impart positional information that governs vertebral identity (9). In mammals, the 39 *Hox* genes are clustered at four separate genomic loci (*HoxA*, *HoxB*, *HoxC*, and *HoxD*), with each gene classified into 1 of 13 paralogous groups dependent on sequence similarities and relative positions within the respective clusters (Fig. 1A). These genes are expressed in partially overlapping domains during embryonic development, with a spatiotemporal collinearity that reflects genomic ordering (10, 11). Exhaustive analysis of *Hox* mouse mutants over more than 20 years has revealed individual and cumulative *Hox* function in conferring specific positional identities to the forming vertebral column (9). For instance, the central/trunk *Hox* proteins (paralogs 5–8) primarily pattern thoracic vertebra, whereas *Hox* 11 paralogs pattern sacral and caudal vertebra (12) and position the sacrum (13, 14).

In addition to transcripts encoding for the *Hox* proteins, transcription within the genomic *Hox* clusters produce noncoding regulatory RNAs, including several microRNAs (miRNAs) (Fig. 1A) (15). In mice, these include the miR-10 family, which is found

Significance

The vertebral column provides essential structural and protective functions. The total number of vertebral elements and their specific morphologies are remarkably reproducible within a given species, yet can be tailored to the requirements of separate vertebrate species. Major genetic determinants driving formation of the vertebral column are known, but how they are regulated to achieve a highly reproducible structure remains to be fully elucidated. In this report, we show that the miR-196 family of microRNAs are essential in defining correct vertebral number and vertebral identity in mouse. We reveal the molecular landscape controlled, either directly or indirectly, by miR-196 activity, to demonstrate that miR-196 impacts many key developmental signalling pathways and reinforces a timely trunk-to-tail *Hox* code transition.

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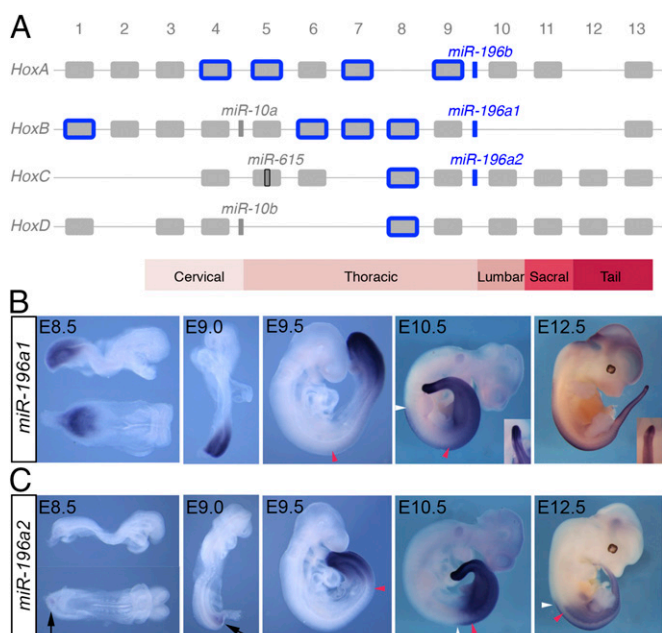


Fig. 1. Unique and overlapping expression patterns of *miR-196* paralogs in mouse. (A) Mouse *Hox* clusters, with the position of *Hox*-embedded microRNAs depicted. Predicted *Hox* targets of the *miR-196* family are indicated in blue. (B and C) Detection of *eGFP* transcripts in *miR-196a1*^{GFP/+} (B) and *miR-196a2*^{GFP/+} (C) embryos demonstrates spatiotemporal expression differences for these identical miRNAs. Embryonic age indicated: red and white arrowheads indicate the anterior boundary of somitic and neural expression, respectively. A discrete band of reduced *eGFP* signal in the anterior PSM of later stage *196a1*^{GFP/+} embryos is shown in *Inset*s. Weak ventral *eGFP* signal in *miR-196a2*^{GFP/+} embryos at E8.5 and E9.0 is indicated with black arrows.

throughout most bilaterian animals; *miR-615*, which is found in eutherian mammals; and the *miR-196* family, which is found in vertebrates and tunicates. Three murine *miR-196* paralogs exist (referred to as *196a1*, *196a2*, and *196b*), each with essentially identical targeting potential (16, 17). The three *miR-196* paralogs exhibit deep conservation across all vertebrate lineages analyzed to date, both in terms of their genomic positioning upstream of *Hox9* paralogs and in their extensive predicted targeting of *Hox* 3' UTRs primarily of the trunk region (Fig. 1A) (17–19). In an early developmental context, *in vivo* validation of these interactions has focused primarily on a single *Hox* target, *Hoxb8* (17, 20–24), with no evidence for additional *Hox* target regulation observed in *miR-196* knockdown studies in zebrafish (24). Thus, the extent to which collective *Hox* output is regulated by *miR-196*, either in terms of the number of genes affected or the relative levels of regulation, is unknown.

The extent to which the developmental modules that define total vertebral number are integrated with those that impart positional information has not been well established, although these processes can be uncoupled (25–27). A function for *Hox* proteins in establishing total vertebral number has been largely dismissed because, with the exception of *Hoxb13*^{-/-} (28), *Hox* knockouts (KOs) do not phenotypically support such a role (9). However, ectopic trunk *Hox* activity can, under certain conditions, drive axis elongation (29). Conversely, posterior *Hox* activity slows axis elongation and terminates the main body axis (29, 30), suggesting an alternative view of *Hox* activity in this context. In this light, phenotypic observations following reduced activity of *miR-196*, a repressor of *Hox* activity, are quite remarkable. Knockdown studies in chick and zebrafish support a role for *miR-196* in regulating vertebral identity (22, 24). Additionally, *miR-196* morphant zebrafish exhibit an extended vertebral column, with what

appears to be an “insertion” of a rib-bearing precaudal element (24). How this latter phenotype arises developmentally is not known and is difficult to reconcile with derepression of trunk *Hox* target genes alone (31, 32). These knockdown approaches could not shed light on individual paralog contributions for this highly related miRNA family, and importantly, the molecular networks downstream of *miR-196*, which have the potential to drive phenotypic alterations, remain uncharacterized.

Here, we generated individual KO alleles for each of the three *miR-196* family members in mice. These tools have allowed us to build an entire allelic deletion series to reveal the individual and additive roles of *miR-196* paralogs in patterning vertebral identity at many axial levels and in controlling the total number of vertebrae. We characterized the detailed molecular landscape controlled by *miR-196* activity in the early embryo to show that *miR-196* regulates, and therefore has the ability to integrate, multiple key signaling pathways to drive developmental processes.

Results

Differential Transcription of *miR-196a1* and *miR-196a2* in the Developing Embryo.

To reveal the individual expression patterns, and therefore potential for functional redundancy, of identical miRNAs *196a1* and *196a2*, we generated *eGFP* knock-in alleles termed *196a1*^{GFP} and *196a2*^{GFP} (Fig. S1). Expression of reporter mRNA reflects sites of active transcription, although it does not reveal additional posttranscriptional regulation that endogenous miRNAs may undergo. Whole mount *in situ* hybridization analysis of reporter mRNA indicated that both miRNAs were expressed specifically in the posterior embryonic derivatives of all three germ layers and revealed striking differences in their spatiotemporal kinetics that have not previously been delineated (Fig. 1B and C). *miR-196a1* is expressed throughout the posterior growth zone at embryonic day (E)8.5 (Fig. 1B). Strong expression is maintained in the PSM until the end of axis elongation, with a discrete band of low expression in the anterior PSM from E10.5 (Fig. 1B, *Inset*). The anterior boundary of somitic and neural expression extends to approximately somite 13/14 [prevertebra (pv) 9, thoracic (T) 2] at E9.5 with a caudal shift in somitic tissue and a rostral shift in neural tissue as development proceeds. This expression profile indicates that *196a1* exhibits a classic collinear profile relative to the adjacent *Hox* gene, *Hoxb9* (anterior limit at E9.5, pv3) (33). *miR-196a2* expression is temporally delayed relative to *196a1* (Fig. 1C), with faint expression ventral to the PSM at E8.5–9.0. Strong expression is then observed throughout the PSM and neural plate at E9.5. A stable anterior somitic limit at approximately somite 21/22 (pv17, T10) and neural limit 2 somites rostral to this is established soon after, consistent with its positioning between *Hoxc9* and *Hoxc10* (34). This analysis revealed both unique and overlapping expression patterns of *196a1* and *196a2*, suggesting these identical miRNAs might have both unique functions where individually expressed and either redundant or additive functions at sites of coexpression.

Genetic Deletion of *miR-196* Leads to Altered Vertebral Identity.

The collective function of *miR-196* family members has yet to be assessed in mammals. Moreover, the dissection of paralog contributions to overall *miR-196* activity has not been achieved in any system. To address this, we generated straight KO alleles at each of the three murine *miR-196* loci (Fig. S2), allowing us to create the complete allelic series of single, double, and triple *miR-196* KO embryos. Skeletal analysis across this allelic series allowed us to demonstrate an essential requirement for *miR-196* activity in patterning the midthoracic, the thoraco-lumbar transition, and lumbo-sacral regions, with both paralog-specific and additive effects revealed.

Removal of individual *miR-196* paralogs alone revealed partially penetrant homeotic patterning defects (Fig. 2A and Table S1). In *196a2* or *196b* single-mutant embryos, the presence of an ectopic rudimentary rib nubbin on the first lumbar vertebra indicated an anterior homeotic transformation of this element (Fig. 2A).

Additionally, in approximately one quarter of cases, we observed anterior homeotic transformations encompassing all subsequent lumbar and sacral elements, resulting in a posterior displacement of the sacrum (schematized in Fig. 2D). Although this latter phenotype could be interpreted as an insertion of a thoracic element, the repositioned last lumbar vertebrae (L6* in Fig. 2D) was often asymmetric, with both lumbar and sacral characteristics

(Table S1), which supports the interpretation of serial identity changes, beginning at L1 and encompassing all subsequent elements. We did not observe a similar L1-to-T anterior homeotic transformation in *196a1* single-mutant embryos, which for the most part exhibited no overt vertebral alterations (Fig. 2A). However, at very low penetrance (Table S1), *196a1* single-mutant embryos displayed an anterior displacement of the sacrum, with or without a reduction

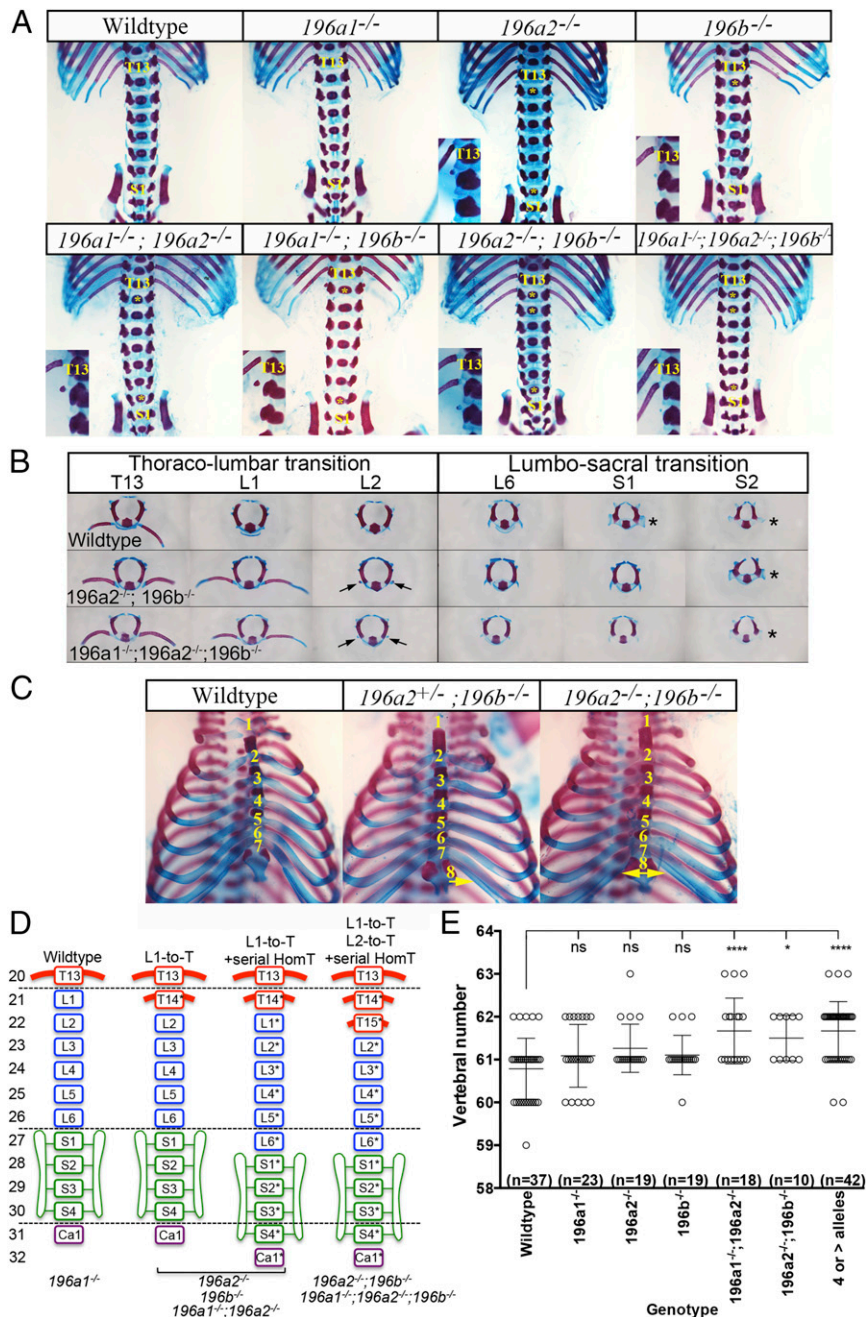


Fig. 2. miR-196 paralogs function in establishing vertebral identity and number in mouse. (A) Identification of vertebral patterning defects in individual and compound *miR-196* loss-of-function E18.5 embryos. Genotypes indicated. The positions of the 13th thoracic element (T13) and first sacral element (S1) are labeled. (Inset) Thoraco-lumbar junction. (B) Individual vertebra analysis to demonstrate identity alterations at the thoraco-lumbar and lumbo-sacral junctions. Genotypes indicated. The position of a rib-like nubbin on lumbar elements is marked with arrow. The position of sacral process is marked with an asterisk. (C) Rib fusion defects observed following loss of *miR-196* alleles, genotypes indicated. Fusion of the eighth rib to the sternum was unilateral or bilateral as indicated with arrows. (D) Summary of patterning defects identified across the *miR-196* allelic series. An asterisk indicates a homeotic transformation of that vertebral element. (E) Quantification of vertebral number in single and compound *miR-196* loss-of-function E18.5 embryos identifies a role for *miR-196* in controlling axis length in mouse. Statistical comparison of vertebral number relative to WT was performed using a permutation test, with *P* values corrected for multiple hypothesis testing using the Bonferroni method: **P* < 0.05, ****P* < 0.001, and *****P* < 0.0001.

in rib length of the last thoracic element (T13), suggesting these paralogs may have an opposing role at this axial level.

We hypothesized that the penetrance and severity of the phenotypes observed after mutating single *miR-196* paralogs could be enhanced by combining these mutations. Indeed, *196a2^{-/-};196b^{-/-}* double-mutant skeletons exhibited a fully penetrant phenotype, with two pairs of supernumerary ribs and anterior homeotic transformation of all subsequent elements (Fig. 2 *A, B*, and *D*). Relative to this double mutant phenotype, triple KO embryos, *196a1^{-/-};196a2^{-/-};196b^{-/-}*, displayed no additional patterning defects (Fig. 2 *A* and *D*, complete dataset summarized in Fig. S3).

We also hypothesized that combining these mutations might reveal additional defects not observed in single mutants. Indeed, all double-mutant skeletons, or skeletons with a triple KO combination of four or more alleles removed, exhibited a partially penetrant increase in the number of ribs attached to the sternum (Table S1 and Fig. 2C), indicating a transformation of the eighth thoracic element to a more anterior identity. Together, our analysis showed that (i) *196a2* and *196b* have single and additive effects in patterning the thoraco-lumbar transition and in positioning the sacrum, with a possible opposing role or *196a1* at this axial level, and (ii) *196a1*, *196a2*, and *196b* act redundantly to pattern the midthoracic region, with phenotypic alterations observed only when two or more paralogs are removed. As such, our work has provided the first genetic proof, to our knowledge, for *miR-196* as a homeotic family of genes and revealed identity changes at multiple axial levels.

Genetic Deletion of *miR-196* Leads to an Increase in Vertebral Number. Homeotic transformations do not alter the number of vertebrae, simply their identity. It was therefore surprising that,

in zebrafish, *miR-196* has been shown to constrain total vertebral number (24). We assessed whether this was an evolutionarily conserved function of *miR-196* and found that the three murine *miR-196* paralogs constrain total vertebral number in a redundant fashion. Wild-type (WT) C57BL6J/N mice exhibit small variations in the total number of vertebrae (Fig. 2E). Compared with the WT mean, we observed a statistically significant increase of approximately one vertebral element in various allelic combinations, including *196a1^{-/-};196a2^{-/-}*, *196a2^{-/-};196b^{-/-}*, and triple KO combinations with four or more alleles deleted (Fig. 2E). Depending on the exact allelic combination, this additional element was patterned as a thoracic (e.g., in *196a2^{-/-};196b^{-/-}* mice) or a postsacral (e.g., in *196a1^{-/-};196a2^{-/-}* mice) element. Together, these results indicate that *miR-196*-mediated control of vertebral number and patterning of segment identity are separable processes. All three *miR-196* paralogs contribute additively to establishing vertebral number within mouse. Layered on top of this, individual *miR-196* paralogs have a differential impact on positional identity and ultimate axial formulae, likely as a result of their differential spatiotemporal kinetics (Fig. 1 *B–K*) relative to target mRNAs.

Transcriptome Alterations Are Detected Following Allelic Removal of *miR-196* Activity. To elucidate the molecular mechanism and targets downstream of *miR-196*, we examined the response of mRNAs to the loss of *miR-196* alleles in E9.5 embryos. To focus these molecular analyses on the relevant cells, i.e., those cells that normally express *miR-196*, we used only embryos with at least one *eGFP* knock-in allele and performed RNA-seq on RNA isolated from cells that were GFP positive (Fig. 3A). With mRNA profiled across 10 genotypes (Table S2), we then compared mRNA

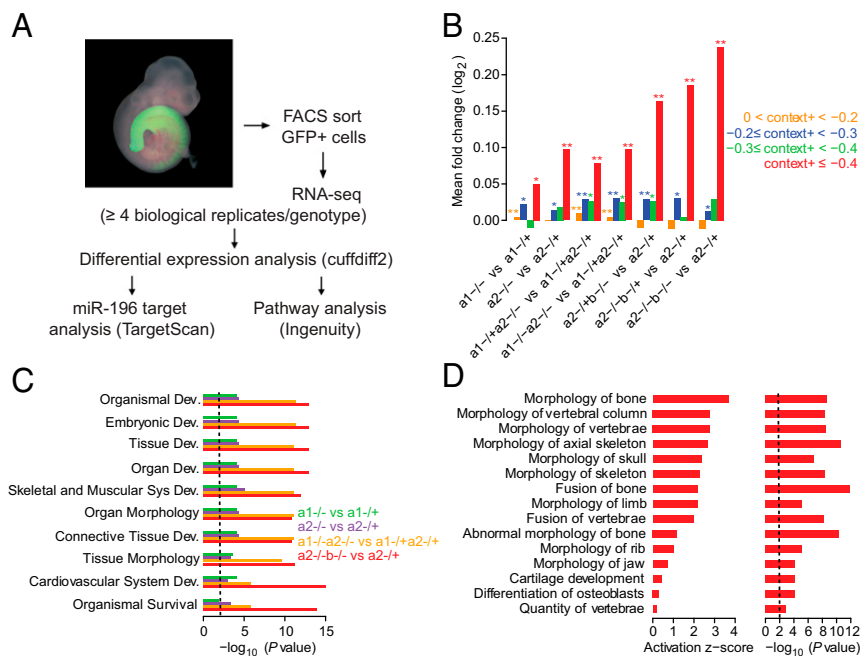


Fig. 3. Whole transcriptome analysis of *miR-196* mutant cells reveals a dysregulation of miRNA targets and skeletal genes. (A) Overview of the experimental and computational strategy used to identify global transcriptome alterations following loss of *miR-196* function. (B) Mean fold changes of genes associated with predicted targets of *miR-196*, partitioned into four context+ intervals according to predicted miRNA targeting efficacy (0 < context+ < -0.2, $n = 2,112$; -0.2 ≤ context+ < -0.3, $n = 145$; -0.3 ≤ context+ < -0.4, $n = 50$; context+ ≤ -0.4, $n = 37$), across seven genotype comparisons. Statistical comparison of observed up-regulation of genes relative to genes with no miRNA target site, as evaluated by a one-sided Kolmogorov–Smirnov (K-S) test; * $P < 0.05$, ** $P < 0.001$. (C) Top 10 significant categories related to gene development and function associated with differentially expressed genes. (D) Top 15 categories related to skeletal and muscular development activated in the *196a2^{-/-};196b^{-/-}* vs. *196a2^{+/+}* comparison, with corresponding activation z-scores and P values. An activation z-score is a measurement of the consistency between the observed pattern of up- and down-regulation of genes in a category and the predicted activation or inhibition pattern in networks stored in the Ingenuity Knowledgebase relative to a random pattern (83). P values in C and D are Benjamini–Hochberg corrected P values, with dashed black lines indicating a significance threshold of 0.01.

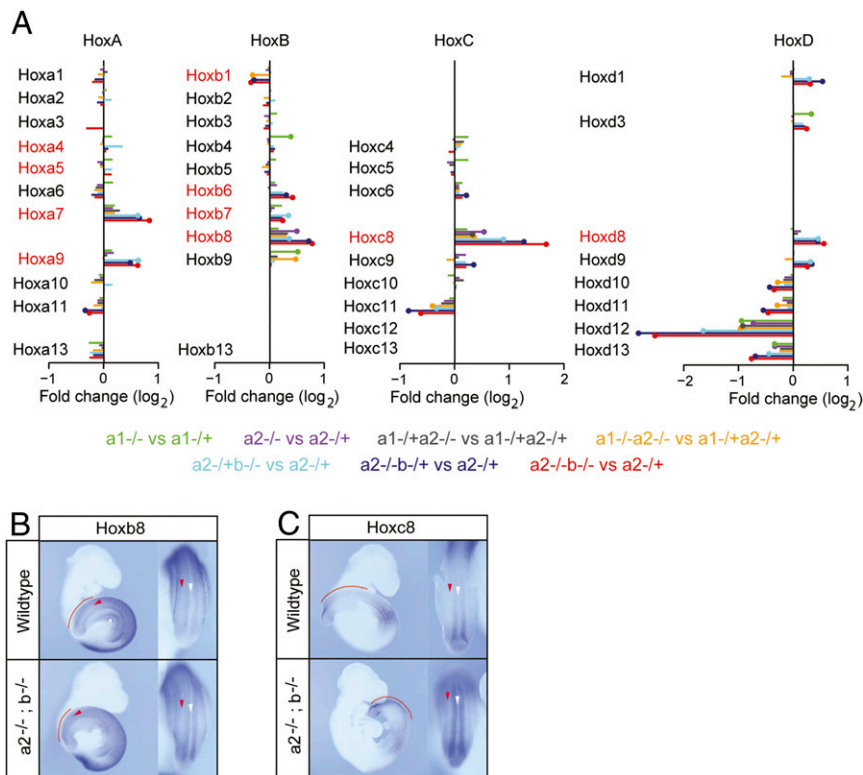


Fig. 4. Loss of *miR-196* function alters global *Hox* signatures. (A) Extensive *Hox* gene dysregulation is identified following loss of *miR-196*. Quantitative expression analysis of all 39 *Hox* genes in cells isolated from E9.5 mutant embryos; genotype comparisons are color coded. *Hox* genes with one or more predicted *miR-196* target binding sites are indicated in red. Filled circles at the tips of fold changes represent a statistically significant change at $q < 0.05$. (B and C) WISH analysis of *miR-196a2^{GFPiGFP};miR-196b^{-/-}* E9.5 embryos relative to WT identifies a caudal expansion of *Hoxb8* (B; $n = 3/3$) and *Hoxc8* (C; $n = 3/3$). The PSM is indicated with a red line/arrowhead and neural tube with a white arrowhead.

changes as increasing numbers and differing combinations of alleles were deleted (Dataset S1). We first examined the effect of allelic *miR-196* deletion on predicted *miR-196* target genes. Using the total context+ score from TargetScan 6.2, which considers the number and type of miRNA binding sites, as well as additional features to predict the genes most effectively targeted by each miRNA (35), we observed that the top predicted targets of *miR-196* exhibited significant derepression on the loss of additional *miR-196* alleles (Fig. 3B and Fig. S4). The derepression of these predicted targets increased with the number of additional alleles deleted (Fig. 3B), revealing *miR-196* dosage sensitivity. The direct interaction between *miR-196* and its target transcripts could occur in any of the three germ layer derivatives in which *miR-196* was expressed, and indeed, an unbiased analysis of all differentially expressed genes revealed statistically altered molecular signatures reflecting this (Fig. S5). Of particular interest, we observe statistical enrichment in genes controlling skeletal morphology (Fig. 3C and D and Fig. S5), indicating the presence of a molecular signature consistent with the vertebral abnormalities observed at the phenotypic level.

Hox Cluster Expression Dynamics Are Altered in *miR-196* Mutant Embryos. It was not known exactly how many of the 10 predicted murine *miR-196* *Hox* target genes are in fact bona fide targets in an in vivo developmental context, nor was it known the relative level of regulation that these predicted targets undergo. When specifically interrogating our transcriptome datasets to assess effects on *Hox* gene expression, a significant and dose-dependent up-regulation of predicted *miR-196* *Hox* targets was observed (Fig. 4A), which paralleled the dose-dependent patterning defects (Fig. 2A). Comparison of *196a2^{-/-};196b^{-/-}* vs. *196a2^{+/-}* profiles

identified 7 of 10 predicted *miR-196* *Hox* targets as significantly derepressed in double-mutant cells at this developmental stage. Those predicted *Hox* targets exhibiting no significant derepression in our analysis included *Hoxb1*, *Hoxa4*, and *Hoxa5*. The most highly derepressed *Hox* targets were *Hoxc8* and *Hoxa7*, both of which harbor multiple predicted *miR-196* binding sites in their 3' UTRs, and *Hoxb8*, which exhibits unusually extensive complementarity to *miR-196* (17, 23). Further, the measurement of differential expression (Fig. 4A) was almost certain to be an underestimate, because our strategy used *eGFP*-positive control samples in which at least one *miR-196* allele had been removed. Whole mount in situ hybridization (WISH) further revealed that the derepression of *Hoxb8* and *Hoxc8* target transcripts in *196a2^{-/-};196b^{-/-}* E9.5 embryos relative to WT manifested as a posterior expansion of endogenous expression domains in both the PSM and neural tube (Fig. 4B and C; $n = 3/3$ per genotype, respectively). In light of previous reports (31, 32), this failure in timely clearance of the trunk *Hox* program from more posterior locations is likely to drive supernumerary rib formation observed in *miR-196* mutant embryos.

Importantly, we also identified a dose-dependent down-regulation of posterior *Hox* genes following progressive removal of *miR-196* alleles (Fig. 4A). This down-regulation was particularly evident for *Hoxd10-d13* genes and was also significant for posterior genes of the *HoxA* and *HoxC* clusters. To confirm that these molecular alterations indeed represented a delay in the activation of a posterior *Hox* program, rather than an overall developmental delay in mutant embryos, we performed quantitative PCR analysis of 17 *Hox* genes in somite-matched WT and *196a2;196b* mutant embryos (somite range, 20–25). This analysis revealed an altered *Hox* signature in mutant embryos (Fig. S6) that was remarkably consistent with that identified by RNA-seq (Fig. 4A). Although the

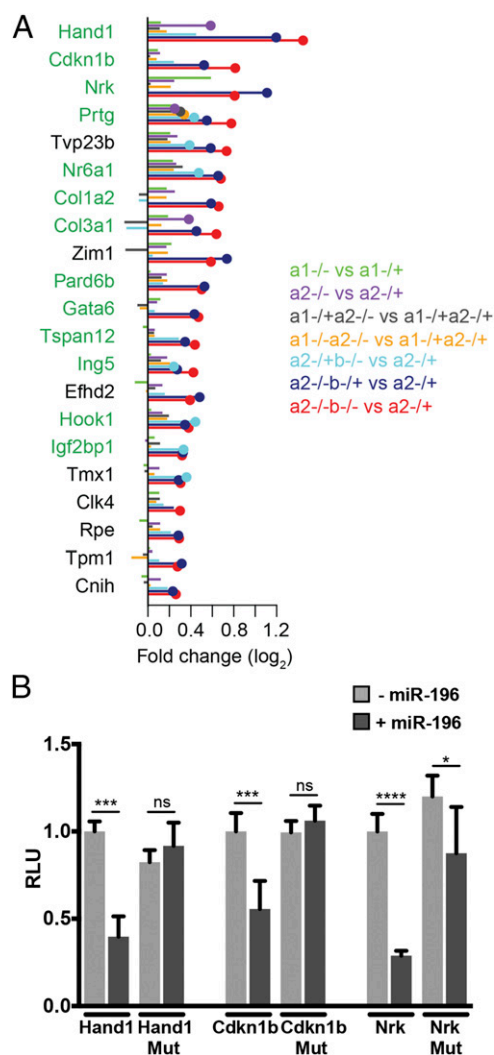


Fig. 5. Identification of additional putative direct (non-*Hox*) miR-196 targets. (A) List of the most highly up-regulated genes and their associated fold changes in seven genotype comparisons that either (i) contain a conserved *miR-196* binding site or (ii) are predicted to respond strongly to the miRNA (i.e., have a context+ score ≤ -0.2). Genes with one or more conserved *miR-196* target binding sites are indicated in green. (B) In vitro luciferase analysis confirms sequence-specific regulation of three experimentally supported target genes of miR-196. Renilla luciferase intensity values have been normalized to their respective Firefly values (RLU). Controls (WT 3' UTR construct without miR-196b) were set to 1. MUT, mutated 3' UTR construct destroying miR-196 binding site. Error bars represent SD. *P* values, Student *t* test: **P* < 0.05, ****P* < 0.0005, and *****P* < 0.0001.

absence of predicted miR-196 sites within these posterior *Hox* mRNAs, together with the direction of the regulation (down instead of up with diminished miRNA), indicated that this regulation was indirect, it was nonetheless notable for three reasons. First, given the potential for phenotypic dominance of posterior over anterior *Hox* protein output (e.g., rib suppression role of *Hox10* paralogs (12, 36), a timely activation of a posterior developmental program in *miR-196* mutants would be expected to suppress supernumerary rib formation. Second, these posterior *Hox* proteins, particularly *Hoxd11* and *Hoxa11*, are known to position the lumbo-sacral junction (13, 37, 38), providing a molecular explanation for how the sacrum was repositioned in *miR-196* mutants. Finally, in addition to understanding vertebral identity defects, these molecular alterations may provide important experimental support for a proposed model whereby maintenance of tailbud cell

divisions, and therefore total vertebral number, is promoted by trunk *Hox* proteins and antagonized by caudal *Hox* proteins (28–30). Our results place *miR-196* activity at this critical junction, coordinating a reproducible trunk-to-tail *Hox* code transition. We suggest that a delay in *Hox* code transition could contribute to the formation of an additional vertebral element observed following genetic removal of miR-196 activity in mice. Such a role for miR-196 is likely to be broadly conserved across vertebrate species, as supported by regionalized vertebral expansion observed in *miR-196* morphant zebrafish (24).

Identification of Additional Direct Targets of miR-196. The statistical enrichment of *Hox* genes among all miR-196 predicted targets (18) prioritized these mRNAs for immediate analysis. However, microRNAs can simultaneously repress extensive suites of target genes (16). To provide experimental support for additional direct targets of miR-196 that have the potential to function in this developmental context, we identified the most highly up-regulated genes in our RNA-seq dataset that either contained a conserved binding site or were predicted to respond strongly to the miRNA (i.e., context+ score ≤ -0.2 ; Fig. 5A). For the top three evolutionarily conserved miR-196 target genes identified, we assessed whether regulation of their expression by miR-196 required direct binding to sites within their 3' UTR. Using a luciferase-based reporter assay system in cell culture, miR-196 was shown to repress each of the target genes in a sequence-specific manner (Fig. 5B). Of particular interest within this set was the cell adhesion molecule (*Prtg*) involved in the ingression of PSM progenitors (39) and an orphan nuclear receptor (*Nr6a1*) essential for somitogenesis in mice (40) and one of the very few genes that has been associated with variation of vertebral number (41). These experimentally supported miR-196 targets highlight important avenues for future investigation, not only with respect to axial patterning and elongation but also the many other

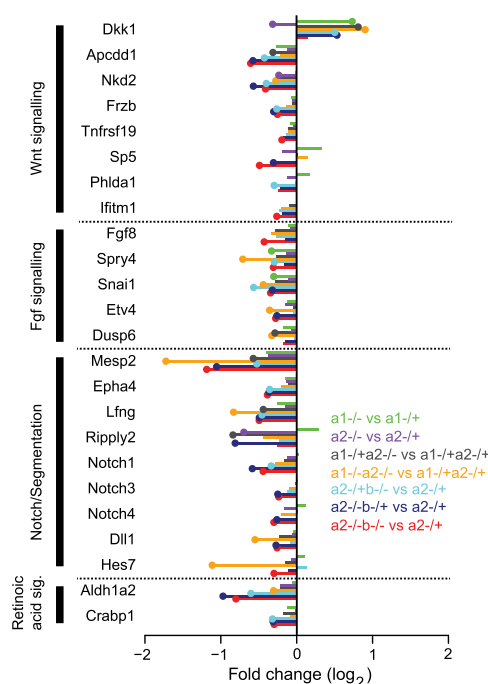


Fig. 6. Loss of *miR-196* function alters signaling pathways known to control segmentation and axis elongation. Quantitative expression analysis of pathways known to control segmentation and axial extension in cells isolated from E9.5 mutant embryos, genotype comparisons are color-coded. Filled circles at the tips of fold changes represent a statistically significant change at *q* < 0.05.

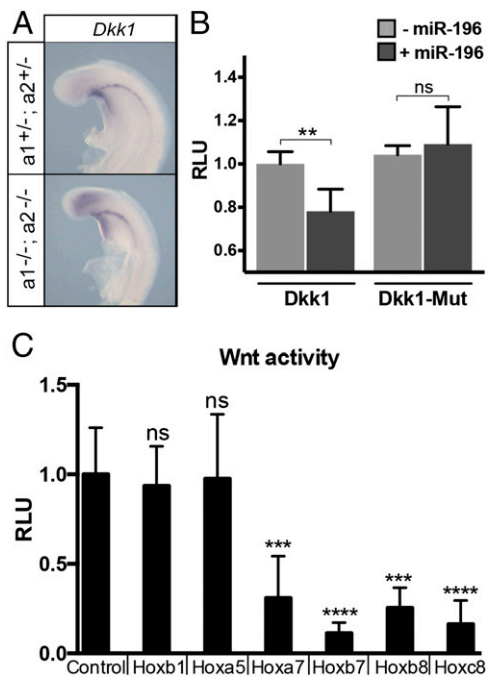


Fig. 7. miR-196 has the potential to regulate Wnt signaling by both direct and indirect mechanisms. (A) WISH analysis confirms increased *Dkk1* in *196a1*^{-/-}; *196a2*^{+/-} E9.5 embryos relative to *196a1*^{-/-}; *196a2*^{-/-} ($n = 2/2$ for each genotype). (B) In vitro luciferase assay confirms sequence-specific regulation of *Dkk1* by miR-196. Renilla luciferase intensity values have been normalized to their respective Firefly values (RLU). Controls (WT 3' UTR construct without miR-196b) were set to 1. MUT, mutated 3' UTR construct destroying miR-196 binding site. (C) Luciferase assay measuring Wnt/ β -catenin activity after overexpression of BATLuc together with CMV-Renilla and either control, *Hoxb1*, *Hoxa5*, *Hoxa7*, *Hoxb7*, *Hoxb8*, or *Hoxc8*; $n = 4-9$ samples per gene assessed. Firefly luciferase intensity values have been normalized to their respective Renilla values (RLU). Control values were set to 1. In B and C, error bars represent SD. Reported P values are from the Student t test: * $P < 0.05$, ** $P < 0.005$, *** $P < 0.0005$, and **** $P < 0.0001$.

developmental processes (20, 21, 24) and pathological conditions (42, 43) involving miR-196.

miR-196 Activity Is Required for Signaling Pathways Associated with Axis Elongation, Segmentation, and the Trunk-to-Tail Transition. miR-196 activity has been shown to negatively regulate retinoic acid pathway activity in the context of pectoral fin formation (24), but regulation of additional developmental signaling pathways in the early embryo, either directly or indirectly, has not been systematically assessed. On further interrogation of our RNA-seq data, we found altered molecular signatures of both axis elongation and somite segmentation across many allelic comparisons (Fig. 6). We observed a clear up-regulation of the Wnt negative feedback inhibitor *Dkk1* (44). In addition, the collective down-regulation of numerous direct and indirect downstream targets of Wnt signaling (45–49) (Fig. 6), and the prediction of diminished β -catenin/*CTNNT1* activity following global pathway analysis (Fig. S7), indicated an overall reduction in Wnt activity in mutant embryos. Wnt and Fgf signaling positively reinforce one another in the mouse tailbud (50–52), and consistent with diminished Wnt activity in *miR-196* mutants, we also observed a down-regulation of the *Fgf8* ligand and numerous Fgf downstream effectors (Fig. 6). We observed a robust down-regulation of Notch signaling components and anterior PSM genes *Mesp2*, *Epha4*, and *Ripply2*, likely as a consequence of diminished Wnt activity acting via the Notch ligand *Dll1* (52–54). Interestingly, these molecular alterations described for *miR-196* mutant embryos resembled alterations

observed following removal of all mature miRNAs in the mesoderm lineage (55), which in the latter case resulted in a caudal displacement of the hindlimb by three somites.

Finally, a coordinated temporal delay in the trunk-to-tail *Hox* code transition has been observed in mice null for *Gdf11* (56), which as heterozygotes, bear striking phenotypic resemblance to *196a2*^{-/-}; *196b*^{-/-} or *miR-196* triple KO mouse embryos. We therefore specifically interrogated our RNA-seq data to assess the levels of *Gdf11* and its direct downstream effector *Isl1* (57). In *196a2*^{-/-}; *196b*^{-/-} embryos, which exhibit 100% penetrant L-to-T transformation and sacral displacement, we observed a statistically significant reduction in *Gdf11* and *Isl1* levels by 15% (Dataset S1). As mentioned, this is likely to be an underestimate of the level of regulation, given the experimental strategy used. The requirement for *Gdf11* in defining presacral vertebral number is dose dependent (56). The exact threshold requirement for *Gdf11* signaling is not known, and it remains to be determined whether subtle down-regulation of *Gdf11* contributes to phenotypic alterations observed in *miR-196* mutant mice. Together, our transcriptome analyses revealed multiple developmental networks that require miR-196 activity for appropriate control of gene expression and suggest intriguing avenues for future experimental exploration.

miR-196 Has the Potential to Modulate Wnt Signaling by Multiple Mechanisms. Vertebral progenitors in the epiblast and tailbud are sensitive to the levels of Wnt signaling. Genetic removal of the Wnt3a ligand (58), or conversely, ectopic activation of Wnt3a in the epiblast (59), result in severe axis truncation posterior to the forelimb. Wnt3a expression has been shown to decrease as progenitor cells commit to a paraxial mesoderm fate (60, 61), and sustained Wnt activity disrupts somite formation (51) and somite polarity (59), dependent on timing and method of activation. These observations indicate that careful titration of Wnt levels is essential throughout the process of somite formation. Our data suggest that miR-196 activity is required in maintaining precise levels of Wnt activity (Fig. 6). Mechanistically, this could be achieved in at least two ways. First, miR-196 could directly target genes in the Wnt pathway. Specifically, the potent Wnt antagonist *Dkk1* harbors a single predicted miR-196 site within its 3' UTR, and *Dkk1* expression was up-regulated following removal of miR-196 activity (Fig. 6). Using WISH, we confirmed increased expression of *Dkk1* in *196a1*^{-/-}; *196a2*^{+/-} embryos relative to *196a1*^{+/-}; *196a2*^{+/-} (Fig. 7A; $n = 2/2$ per genotype). To test whether miR-196 can act directly to repress *Dkk1*, we used a luciferase-based reporter assay system in cell culture to show that, indeed, miR-196 negatively regulates the *Dkk1* 3' UTR in a sequence-specific manner (Fig. 7B). However, the repression in the reporter assay was more modest than that observed in vivo using RNA-seq (Fig. 6), and *Dkk1* is not a conserved target of miR-196, suggesting that indirect regulation by miR-196 also plays a role. Second, miR-196 control over Wnt activity might work in part via *Hox* intermediates, which have the potential to either activate or repress Wnt signaling (29, 30). We recently showed using chick in vivo electroporation and imaging that the collinear activation of a subset of *Hox9-13* posterior *Hox* genes within paraxial mesoderm progenitors translates into a graded increase in Wnt repression and a slowing down of axis elongation (30). One *Hox* gene that was found to significantly repress Wnt activity using this in vivo luciferase-based Wnt reporter assay was the miR-196 target *Hoxa9*. We therefore went on to test whether additional miR-196 *Hox* targets have the ability to repress Wnt activity in this context. We coelectroporated a Wnt/ β -catenin reporter (BATLuc) and a CMV-Renilla construct in paraxial mesoderm progenitors together with an expression vector containing either Venus or *Hoxb1*, *Hoxa5*, *Hoxa7*, *Hoxb7*, *Hoxb8*, and *Hoxc8*. Of these six *Hox* genes tested, four (*Hoxa7*, *Hoxb7*, *Hoxb8*, and *Hoxc8*) showed strong repression of luciferase activity, whereas two (*Hoxb1* and *Hoxa5*) did not (Fig. 7C). Interestingly, the two

Hox genes do not influence Wnt/ β -catenin reporter activity in early chick paraxial mesoderm progenitors are the same *Hox* genes that show no indication of direct regulation by miR-196 in E9.5 mouse tissue (Fig. 4A). Together, these data demonstrate that miR-196 has the potential to directly and indirectly regulate the precise levels of Wnt activity in the developing embryo.

Discussion

Our work demonstrates the essential role for murine miR-196 in regulating vertebral identity across different levels of the body axis and reveals evolutionary conservation in the role of miR-196 in constraining total vertebral number. Importantly, our strategy has allowed us to comprehensively dissect paralog contribution to resultant phenotypes, allowing us to distinguish a patterning role for miR-196 from its role in modulating vertebral number. Moreover, we characterized the detailed molecular landscape controlled by miR-196 activity in the early embryo to show that miR-196 regulates, and therefore has the ability to integrate, multiple key signaling pathways to drive developmental processes.

miR-196 Activity Is Essential for Vertebral Identity. Despite the clear potential for functional redundancy between miR-196 paralogs (17), homeotic transformation of vertebral elements could be observed at low penetrance following removal of an individual miR-196 paralog (e.g., *196a2*^{-/-} or *196b*^{-/-} single mutants). With increasing loss of miR-196 family members (e.g., *196a2*^{-/-};*196b*^{-/-} double mutants), fully penetrant vertebral phenotypes were observed that were equivalent in severity to many single and compound *Hox* mutants (14, 62). Vertebral identity changes were observed at sites where loss-of-function phenotypes have previously been described for many of the direct *Hox* target genes (62), reinforcing the view that miR-196 acts within endogenous *Hox* domains rather than simply as a fail-safe mechanism to clear an anterior developmental program at more posterior locations (22). Paradoxically, the *196a2*^{-/-};*196b*^{-/-} or triple KO phenotypes are remarkably similar to either *Hoxc8*^{-/-} or *Hoxc8*^{-/-};*Hoxd8*^{-/-} skeletons, with eight ribs attached to the sternum, L1-to-T transformation, and a posterior displacement of the sacrum (62). However, with respect to number of sternal rib attachments and L1-to-T transformation, *Hoxc8* loss-of-function and gain-of-function mutant mice exhibit identical phenotypes (31, 62). These data indicate that exquisite regulation of a quantitative *Hox* code is essential in defining vertebral identity at this axial location. Interestingly, deletion of *Hoxb8* rescues many defects observed in *Hoxc8*-null mice, highlighting that there are aspects of a qualitative *Hox* code that we are yet to understand. Nonetheless, similar to *Hoxc8*, ectopic *Hoxb8* expression results in supernumerary rib formation throughout the lumbar region (32), supporting the view that a collective up-regulation of direct *Hox* target genes drives homeotic alterations of the midthoracic to upper lumbar region in miR-196 mutant mice.

A shift in the position of the sacrum observed in miR-196 mutant embryos was not easily reconcilable with the function of miR-196 in directly repressing trunk *Hox* target genes (31, 32). However, we show that in addition to direct *Hox* gene regulation, miR-196 indirectly regulates the expression levels or temporal activation of many caudal *Hox* genes, including those whose protein products are known to control positioning of the sacrum, such as *Hoxa10*, *Hoxd10*, and *Hoxd11* (13, 14, 63, 64). The mechanisms leading to a delay in posterior *Hox* gene activation in miR-196 mutant mice are currently unknown. A similar coordinated temporal shift in the trunk-to-tail *Hox* code has been demonstrated in *Gdf11*^{-/-} mice (56), which show conservation in the types of vertebral transformations we observe here in miR-196 mutant embryos. In this context, *Gdf11* appears to work via retinoic acid signaling (57, 65), and whether altered *Gdf11* and retinoic acid signaling contribute to miR-196 phenotypic alterations remains to be tested.

miR-196 Activity Constrains Total Vertebral Number. Total vertebral number of a given species is highly reproducible, and mutations that extend the vertebral column of model organisms are very rare. Among vertebrate species, however, great diversity in vertebral number has arisen. Cross-species comparison (8) or direct genetic perturbation (25, 27) demonstrates that the periodicity of segmentation clock oscillation relative to the rate of PSM growth is the central parameter in defining vertebral number. It remains to be determined how an additional vertebral element seen here in miR-196 mutant mice, or in miR-196 morphant zebrafish (24), are generated at a cellular level (i.e., does the clock tick faster or does it tick at the same rate for longer). Our analysis does, however, reveal molecular alterations in miR-196 mutant embryos that have the potential to affect vertebral number.

First, altered expression of Notch, Wnt, and Fgf pathways could alter the periodicity of segment formation (1). However, diminished Wnt and Fgf would be predicted to increase somite size (27, 66–68), which if axis elongation was unaltered, would lead to a reduction in vertebral number. Further work is required to clarify any functional role for miR-196 in the molecular networks coordinating segmentation.

Second, we showed that miR-196 activity can modulate the expression levels of many *Hox* genes, either directly or indirectly. It is well documented that *Hox* genes control mesodermal ingression, thus regulating cell injection into the PSM (30, 69). The rate of PSM growth is not uniform along the anterior-posterior axis (8), with a switch to PSM shortening occurring at about the trunk-to-tail transition in most amniotes. This switch correlates with activation of a posterior *Hox* code (*Hox9* onward), and a subset of posterior *Hox* genes slows axis elongation by controlling the ingression of PM progenitors via Wnt repression (30). We show here that the ability to repress Wnt signaling is not exclusive to posterior *Hox* genes but that *Hox7/8* paralogs also down-regulate Wnt signaling in the chick epiblast. The repression of Wnt by posterior *Hox* genes as a means to slow down and terminate axis elongation (29, 30) is consistent with the known function of Wnt3a in driving axis elongation (58). The repression of Wnt by trunk *Hox* genes is less intuitive because, at least on a *Cdx2/4* mutant background, ectopic *Hoxb8* has been shown to maintain Wnt expression in the tailbud (29). However, the importance of precise Wnt levels in the early steps of axis formation, and of cellular context, are beginning to be appreciated (59, 70) and may underlie such discrepancies. Exactly how miR-196 activity integrates within the genetic networks controlling PM progenitor maintenance and differentiation remains to be fully elucidated. At E9.5, this progenitor population represents quite a minor fraction of miR-196-expressing cells. Therefore, the extent to which molecular alterations we identified in miR-196 mutants, including reduced Wnt/Fgf signaling, reflect an altered PM progenitor molecular status requires clarification at a cellular resolution.

Although a heterochronic shift in the trunk-to-tail *Hox* code transition could be predicted to vary vertebral number, morphological evidence for this has been scarce. Analysis of total vertebral number in *Gdf11*^{-/-} mice, which exhibit a dramatic heterochronic shift in *Hox* code, is hampered by caudal truncation (56). Although ectopic trunk *Hox* expression (*Hoxa5* and *Hoxb8*) has the ability to rescue axis truncation defects of a genetically engineered mutant (29), they do not appear to increase vertebral number on a WT background (29, 31). This lack of phenotype is possibly due to the fact that posterior prevalence still holds; caudal *Hox* genes and miR-196 would be expressed at the usual time and place to regulate and terminate axis elongation. In the case of miR-196 KOs, the cumulative effect on both trunk and caudal *Hox* gene expression could permit continued maintenance of progenitor divisions while delaying commencement of axis elongation slow down, resulting in increased vertebral number.

Together, our results highlight an essential requirement for miR-196 activity in reinforcing a timely trunk-to-tail *Hox* code

transition and reproducibility of axial formulae. Given the ancestral role of Hox activity in species that use a posterior growth zone (71), and the recurrent acquisition of miRNAs within the *Hox* clusters across metazoan taxa (15, 17, 72, 73), variation in *Hox*-miRNA interactions may represent an important mechanism for the evolution of animal body plans.

Materials and Methods

miR-196a1^{GFP} and miR-196a2^{GFP} Knock-In Construction. A 72- (*miR-196a1*) or 52-bp (*miR-196a2*) genomic fragment encompassing each mature miRNA sequence was replaced with a cassette containing eGFP fused to the rabbit β -globin 3' UTR followed by flippase recognition target (FRT)-flanked PGK α -Neomycin. A Kozak sequence was inserted upstream of the eGFP start codon. Targeting constructs were generated using a 129/Sv sequence and electroporated into J1 embryonic stem cells. Correctly targeted ES cells were identified and used to generate germ-line transmitting knock-in lines. Before analysis, the Neomycin selection cassette was removed by crossing to a ubiquitous FLPe-deleter mouse line. Resulting lines were bred onto a C57BL/6J background and confirmed as isogenic by SNP genotyping.

miR-196a1^{-/-}, miR-196a2^{-/-}, and miR-196b^{-/-} Generation. Previously targeted ES cells at each of the three *miR-196* loci have been generated (74). Correctly targeted JM8A3 ES cells were reconfirmed by Southern blot and used to generate germ-line transmitting KO lines. Before analysis, the puDeltaTK selection cassette was removed by crossing to a ubiquitous Cre-deleter mouse. Resulting lines are on a mixed C57BL/6J and C57BL/6N background.

Mouse Skeletal Preparation and Analysis. Skeletal preparation was performed on E18.5 embryos or p0 postnatal pups as previously described (75).

In Situ Hybridization. Whole mount in situ hybridization was performed as previously described (76).

FACS Sorting and RNA-seq Sample Preparation. Freshly dissected E9.5 embryos were dissociated in 0.25% trypsin/2% (vol/vol) chick serum, neutralized in DMEM + 10% (vol/vol) FBS, and washed into PBS + 2% (vol/vol) FBS for FACS sorting. GFP-positive cells were FACS sorted directly into RNeasy lysis buffer (Qiagen) and RNA isolated using RNeasy with added on-column DNase treatment (Qiagen). RNA quality was assessed using a Bioanalyser, and 200 ng per individual embryo was used as input for RNA-seq library generation (unstranded Illumina TruSeq Kit). Libraries were multiplexed and sequenced using an Illumina HiSeq. 2000 instrument, generating 50-bp single end reads.

RNA-seq and Category Enrichment Analysis. Quantification of the transcriptome using RNA-seq data was performed as previously described (77). Raw reads were aligned to the latest build of the mouse genome (mm10) using STAR v. 2.3.1n (options—outFilterType BySJout—outFilterMultimapScoreRange 0—readMatesLengthsIn Equal—outFilterIntronMotifs RemoveNoncanonicalUnannotated—clip3pAdapterSeq TCGTATGCCGCTTCTGCTT—outSAMstrandField intronMotif—outStd SAM) (78). Considering all replicates of a particular genotype, differential expression statistics were computed between genotypes of interest using cuffdiff v. 2.1.1 (options—library-type fr-unstranded -c 100 -b mm10.fa -u-max-bundle-frags 10000000) (79), using mouse transcript models of protein-coding genes annotated in Ensembl release 72. Before all subsequent analyses, we filtered away genes annotated by cuffdiff as “NOTEST” in all genotypes, indicating the genes were too lowly expressed to accurately quantify their abundances. To evaluate functional gene categories that were statistically enriched, we loaded differentially expressed genes (i.e., genes with $Q < 0.05$) into the Core Analysis function of Ingenuity Pathway Analysis software (Ingenuity Systems), testing gene categories related to development and function. All *P* values reported from this analysis were adjusted using the Benjamini–Hochberg method to control the false discovery rate.

miRNA Target Analysis. To identify predicted miRNA targets, the 3' UTR sequences of protein-coding genes were searched to identify 6mer, 7mer-A1, 7mer-m8, and 8mer miRNA binding sites cognate to the miR-196 seed (35, 80). A context+ score was computed for each target site within a given 3' UTR, and scores were summed to produce a total context+ score for each gene, which was used for all miRNA-related analyses (35). TargetScanMouse 6.2 was further used to assess target site conservation or to include predicted miR-196 targets containing noncanonical 3' compensatory sites, such as in the case of *Hoxb8* (81).

Permutation Test for Significance Testing. A permutation test was devised to evaluate the significance of differences in vertebral number. Briefly, given two groups of count-based data of size *n* and *m*, we randomly partitioned the counts (without replacement) from the union of the two groups to generate 100,000 pairs of data, again of size *n* and *m*. To compute an empirical one-sided *P* value, we then computed the proportion of pairs that satisfied the condition that the difference in the means of each pair exceeded the difference in means of the original two groups.

In Vitro Luciferase Assay. A 3' UTR sequence (300–700 nucleotides) of protein-coding genes of interest were commercially synthesized and cloned into psiCheck2 vector. For each, a mutant version containing four nucleotide substitutions within the miR-196 seed sequence was generated. Constructs were transfected into NIH 3T3 cells with or without 25 pmol mmu-miR-196b duplex. Transfection (Lipofectamine2000; Life Technologies) and luciferase analysis (Dual Luciferase Reporter Assay System; Promega) were performed as per the manufacturer's instructions.

Chick Electroporation and in Vivo BatLuc Reporter Analysis. Chicken embryos were harvested at Hamburger and Hamilton stage 5 (82) and electroporated ex ovo as previously described (30), with a DNA mix containing BATLuc (1 μ g/ μ L final), CMV-Renilla (Promega; used as a control to normalize the differences of electroporation intensity between embryos; 0.2 μ g/ μ L final), a control pCAGGS-Venus vector (gift from K. Hadjantonakis, Memorial Sloan Kettering Cancer Center, NY), or a *Hox* gene of interest (*Hoxb1*, *a5*, *a7*, *b7*, *b8*, or *c8*) cloned in pCAGGS-IRES2-Venus (5 μ g/ μ L final). Electroporated embryos were cultured in a humidified incubator at 38 °C for 20 h. Embryos were analyzed using a fluorescent microscope, and only embryos showing restricted expression of Venus in the paraxial mesoderm were selected (90–100% of the electroporated embryos) for luciferase assay (between three and five embryos for each condition). The posterior region (from somite 1 to tailbud) of the selected embryos was dissected and lysed in passive lysis buffer (Promega) for 15 min at room temperature. Lysates were then distributed in a 96-well plate, and luciferase assays were performed using a Centro LB 960 luminometer (Berthold Technology) and the dual luciferase kit (Promega) following manufacturer's instructions. Raw intensity values for Firefly luciferase signal were normalized with corresponding Renilla luciferase values (RLUs), and the control experiment was set to 1.

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