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Multiple inputs into a posterior-specific regulatory network in the *Ciona* notochord

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Abstract

The gene regulatory networks underlying *Ciona* notochord fate specification and differentiation have been extensively investigated, but the regulatory basis for regionalized expression within the notochord is not understood. Here we identify three notochord-expressed genes, *C11.331*, *C12.115* and *C8.891*, with strongly enriched expression in the secondary notochord cells at the posterior tip of the tail. *C11.331* and *C12.115* share a distinctive expression pattern that is highly enriched in the secondary notochord lineage but also graded within that lineage with the strongest expression at the posterior tip. Both genes show similar responses to pharmacological perturbations of Wnt and FGF signaling, consistent with an important role for Wnt and FGF ligands expressed at the tail tip. Reporter analysis indicates that the *C11.331* *cis*-regulatory regions are extensively distributed, with multiple non-overlapping regions conferring posterior notochord-enriched expression. Fine-scale analysis of a minimal *cis*-regulatory module identifies discrete positive and negative elements including a strong silencer. Truncation of the silencer region leads to increased expression in the primary notochord, indicating that *C11.331* expression is influenced by putative regulators of primary versus secondary notochord fate. The minimal CRM contains predicted ETS, GATA, LMX and Myb sites, all of which lead to reduced expression in secondary notochord when mutated. These results show that the posterior-enriched notochord expression of *C11.331* depends on multiple inputs, including Wnt and FGF signals from the tip of the tail, multiple notochord-specific regulators, and yet-to-be identified regulators of regional identity within the notochord.

Introduction

The *Ciona* notochord consists of only 40 cells that intercalate to form a tapered, single-file column. Cell fate specification in the *Ciona* notochord has been extensively studied. In ascidians, the primary notochord cells are derived from blastomere pairs A7.3 and A7.7 and give rise to the anterior 32 cells in the intercalated notochord, whereas the secondary notochord cells are derived from B8.6 and give rise to the posterior 8 cells (Nishida, 1987). In *Ciona*, A7.3 and A7.7 become fate-restricted to notochord at the 64-cell stage downstream of both beta-catenin mediated mesendodermal specification and FGF signaling (Hudson et al., 2016, 2013; Yasuo and Hudson, 2007). B8.6 becomes fate-restricted at the

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112-cell stage via different mechanisms that involve a Nodal/Delta2 signaling relay initiated by the lateral b6.5 cells (Hudson and Yasuo, 2006, 2005, Imai et al., 2006, 2002).

Both primary and secondary notochord cells express the key notochord-specific transcription factor Brachyury (Bra) (Corbo et al., 1997; Yasuo and Satoh, 1993), and the majority of notochord-enriched genes are expressed uniformly throughout the notochord (Hotta et al., 2000, 1999; José-Edwards et al., 2011; Kugler et al., 2008; Reeves et al., 2017). We have previously shown, however, that notochord cell sizes, shapes and behaviors vary depending on both lineage and anterior-posterior position (Carlson et al., 2015; Veeman and Smith, 2013). We have also shown that a subset of notochord-enriched genes are differentially expressed within subregions of the notochord (Reeves et al., 2014). The regulatory basis for regionalized notochord expression has received no previous attention.

Differential gene expression between primary and secondary notochord is of particular interest because secondary notochord cells are quite distinct from primary notochord cells in their morphology and behavior. Secondary notochord cells are more stereotyped in their patterns of intercalation than primary cells and give rise to a particularly distinct taper towards the tail's distal tip (Carlson et al., 2015; Veeman and Smith, 2013). Unlike primary cells, they still intercalate effectively even in mutants for the planar cell polarity pathway gene *Prickle* (Jiang et al., 2005). Elements of differential gene expression have previously been noted between primary and secondary notochord cells, including a gene upregulated at both the anterior and posterior tips of the notochord, one expressed in a posterior to anterior gradient, and another expressed throughout the notochord but with visible enrichment in the secondary lineage (Reeves et al., 2014). Despite these indications of underlying patterning, markers that were strongly differentially expressed between primary and secondary notochord cells were previously elusive.

We recently used RNAseq on flow-sorted notochord cells to identify a large set of 1364 genes predicted to be enriched in the notochord. Extensive *in situ* validation of 151 genes showed that ~90% were indeed notochord enriched (Reeves et al., 2017 and unpublished data). In the course of this we identified several new genes showing regionalized notochord expression, including three that are strongly enriched at the notochord's posterior tip. Here we characterize the expression patterns of these posterior notochord tip-specific genes with fine detail and investigate the molecular mechanisms controlling their regionalized expression.

Results

Secondary-enriched notochord expression

Although the majority of notochord enriched genes we identified by RNAseq are expressed uniformly across the notochord (Reeves et al., 2017), a subset display regionalized expression within the notochord. Fig. 1 shows *in situ* expression patterns for three genes with strongly differential expression between primary and secondary notochord. *C11.331* has homology to the fibulin/hemicentin family of secreted extracellular matrix proteins (Timpl et al., 2003), but is not a clear ortholog of any vertebrate fibulin. *C12.115* and *C8.891* are both predicted SLC family solute carriers. We refer to all three here by their KH2012

gene model names (Brozovic et al., 2018; Satou et al., 2008) for unambiguity. During early notochord intercalation at mid neurula (Hotta stage 16) (Hotta et al., 2007), *C11.331* is barely detectable (Fig. 1A), but is distinctly expressed at the notochord's posterior tip late in intercalation at early tailbud stage 20 (Fig. 1B) and during tail elongation at late tailbud stage 23 (Fig. 1C). *C12.115* has posterior-enriched notochord expression at all three stages (Fig. 1D–F) which resolves over time to be increasingly specific to the notochord's posterior tip. *C8.891* is qualitatively different than the other two genes and shows a more graded expression pattern at the first two stages (Fig. 1G,H) before developing a sharper transition between primary and secondary at stage 23 (Fig. 1I).

To better quantify their expression patterns, we also imaged fluorescent *in situ* hybridizations against *C11.331* and *C12.115* by confocal microscopy. This allowed the boundary between primary and secondary notochord to be clearly identified. Both genes showed expression in secondary but not primary notochord (Fig. 1J,K). For both *C11.331* and *C12.115*, this secondary-specific expression was stronger at the notochord's posterior tip than at the boundary between primary and secondary notochord. Fig. 1L,M show quantitative analyses of these expression patterns across nine embryos each, confirming that both genes have graded expression in the secondary notochord that is strongest at the posterior tip.

Perturbation of Wnt and FGF signaling

These expression patterns suggested the potential involvement of a secreted signaling molecule expressed at the posterior tip of the tail. Candidate molecules for such a signal include Wnt5, FGF8/17/18 and FGF9/16/20, all of which are expressed in tail tip ectoderm or posterior tail muscles (Hudson et al., 2007; Hudson and Yasuo, n.d.; Ikuta et al., 2010; Imai et al., 2004). We tested their involvement pharmacologically using the FGF inhibitor U0126 and the canonical Wnt pathway activator BIO. Both of these drugs have been extensively used in *Ciona* and we used doses previously established to be specific and effective (Dumollard et al., 2013; Hudson et al., 2013; Racioppi et al., 2014; Sakabe et al., 2006). We treated the embryos with drugs starting at mid gastrula (Hotta stage 12). This is after notochord has been specified and has completed all cell divisions (Hotta et al., 2007), but before the earliest detectable expression of *C11.331*, *C12.115* or *C8.891* by *in situ* hybridization (data not shown). A Wnt pathway inhibitor would be conceptually appealing but no such reagents have been validated in *Ciona* embryos.

Both drugs caused mild defects in tail morphogenesis, as has been described previously for U0126 (Ikuta et al., 2010). As expected, the BIO and U0126 treatments had no effect on the expression of the key notochord transcriptional regulator *Brachyury* (Fig. 2A–D), but the drugs caused strong effects on the expression of the three posterior-specific notochord markers. *C11.331* and *C12.115* responded to both drugs in similar ways. BIO treatment led to the expansion of *C11.331* and *C12.115* expression throughout primary notochord (compare Fig. 2E and I to Fig. 2F and J). Interestingly, this ectopic expression was often strongest at the notochord's anterior tip, giving rise to an overall expression pattern reminiscent of *BCamL* (*KH.C2.209*) (Reeves et al., 2014) in terms of being enriched at both the anterior and posterior tips of the notochord. FGF inhibition with U0126, however, led

to a major decrease in the expression of both markers (Fig. 2G,K). This was particularly evident for *C12.115* expression, which was virtually eliminated by U0126 treatment despite being much more robustly expressed than *C11.331* in untreated embryos. *C8.891* expression expanded throughout the notochord in response to Wnt pathway activation by BIO, but this ectopic expression was uniform and did not show the ‘both tips’ pattern usually seen with the other two genes (Fig. 2N). *C8.891* also differed from *C11.331* and *C12.115* in that FGF inhibition by U0126 only caused a modest decrease in expression (Fig. 2O).

The BIO and U0126 treatments indicated that both Wnt and FGF signals may act as positive regulators of *C11.331*, *C12.115* and *C8.891* expression. As a preliminary test of the potential relationships between Wnt and FGF signaling in this context, we also treated embryos with both drugs simultaneously. Both *C11.331* and *C12.115* showed an intermediate phenotype in which expression expanded throughout the notochord but was much fainter than in response to BIO treatment alone (Fig. 2H,L). In addition, no enrichment in the anterior tip was seen and posterior enrichment was also lost in most embryos. This lack of a clear epistatic relationship suggests that Wnt and FGF signals act in parallel upon these two genes. *C8.891* again behaved differently; treatment with combined BIO and U0126 led to expanded expression throughout the notochord, but this ectopic expression was strong and indistinguishable from BIO treatment alone (Fig. 2P). FGF signaling appears to play a less important role in the regulation of *C8.891* compared to the other two posterior enriched genes.

Quantitative *cis*-regulatory analysis

To narrow in on the molecular mechanisms responsible for posterior-specific notochord expression, we developed a dual reporter strategy. Electroporated transgenes in *Ciona* are expressed mosaically, and it is common for individual embryos electroporated with reporter constructs for uniformly expressed notochord genes such as *Brachyury* to exhibit expression in secondary notochord but not primary and *vice-versa* (Carlson et al., 2015; Corbo et al., 1997). Different transgenes electroporated at the same time, however, typically show common patterns of mosaicism (Zeller et al., 2006). To control for differential reporter expression due to transgene mosaicism, as opposed to *bona fide* differences in regionalized expression, we co-electroporated HA-tagged Histone H2B under the control of the *Bra* enhancer/promoter together with Venus fluorescent protein downstream of candidate regulatory regions and a basal promoter (Fig. 3A).

We used this dual reporter approach to investigate the *cis*-regulation of *C11.331*. All embryos were imaged by confocal microscopy using uniform imaging parameters. We developed a quantitative analysis approach based on measuring Venus and HA intensity along the AP axis of computationally straightened and flattened notochords. For each embryo analyzed, we measured the mean background-corrected Venus reporter signal for *Bra*>H2B:HA expressing cells in primary notochord (cells 1–32) and secondary notochord (cells 33–40), and normalized the results to the *Bra*>H2B:HA intensity in each region (Fig. 3B). We also scored a larger number of embryos with a qualitative system, ranging from 0 (no detectable expression) to 4 (oversaturated) (Tables S1,S2), which largely matched the quantitative analysis. An ectopic expression score was generated based on qualitative

expression levels in several non-notochord tissues (Tables S3,S4). Full details of the scoring systems are given in the Methods section.

C11.331 is separated from the 3' end of the closest upstream gene by ~1.5kb, and we found that this intergenic region (-1565 to +13) gave rise to strong expression in secondary notochord, much weaker expression in primary notochord, and relatively little ectopic expression (Fig. 3B). We dissected this region with a series of truncations and found evidence for extensive functional redundancy. On a coarse scale, both the distal -1565 to -758 and the proximal -787 to +13 region gave rise to secondary-enriched notochord expression, with the distal region showing somewhat less ectopic expression but the proximal region showing stronger notochord expression and greater statistical evidence for enrichment in the secondary notochord. We further dissected the -787 to +13 region and again found aspects of redundancy, with both the -787 to -405 and the minimally overlapping -488 to +13 region showing secondary-enriched notochord expression.

Finer-scale analysis of the -488 to +13 region allowed us to identify distinct regulatory regions. A construct spanning the -488 to -165 region was still strongly expressed in secondary notochord. Unlike the previous constructs, however, it was strongly expressed in primary notochord as well, indicating the presence of a silencer element for primary notochord expression proximal to -165. A further truncation to give the -488 to -245 construct eliminated expression altogether, indicating the presence of essential enhancer elements in the -245 to -165 interval. Truncating from the other side, the -322 to +13 construct was highly expressed and strongly enriched in secondary notochord, whereas the -245 to +13 construct was enriched in secondary notochord, but much more weakly expressed overall, indicating that there are additional enhancer elements between -322 and -245. Our subsequent efforts therefore focused on the -322 to +13 interval.

Fig. 4A shows normalized *C11.331*(-322 to +13)>Venus expression as a function of AP position for 19 imaged notochords. The normalized mean intensity (green line) reveals a sharp transition from low to high expression precisely at the boundary between primary and secondary notochord. This is in contrast to the *Bra*>H2B:HA control that is expressed uniformly along the notochord's AP axis (Fig. 4B).

For these finer-scale dissections, we quantified reporter expression separately for the anterior (cells 33–36) and posterior (cells 37–40) secondary notochord. For the -322 to +13 construct, the sample mean of this quantitative metric of reporter expression was 19.7 fold higher in anterior secondary notochord as compared to primary notochord, and 26.6 fold higher in posterior secondary notochord as compared to primary notochord. These differences were highly significant as measured by the Wilcoxon sign-rank test. Two truncated constructs, -322 to -92 and -322 to -127, retained significant enrichment in secondary notochord but were less highly expressed overall, indicating that there are enhancer elements promoter-proximal to -92. A further truncation to give the -322 to -165 construct was significantly derepressed in primary notochord, confirming the presence of important silencer elements promoter-proximal to -165. Combined with the previous constructs (Fig. 3B), this indicates that the primary notochord silencer has essential elements between -165 and -127.

Identification and testing of candidate TFBSs

We searched the –165 to –127 interval and flanking regions for putative transcription factor binding sites (TFBSs) using both TRANSFAC position weight matrices (PWMs) (Matys et al., 2006) via the LASAGNA web tool (Lee and Huang, 2013) and also using SELEX-seq-derived (Slattery et al., 2011) consensus sequences for known *Ciona* notochord TFs downloaded from the ANISEED community database (Brozovic et al., 2018). We found several predicted sites of interest, including an ETS site just outside of the silencer in a region required for strong expression (Fig. 5A). This was of particular interest given that ETS family transcription factors are key mediators of ascidian FGF signaling (Gainous et al., 2015; Miya and Nishida, 2003). We did not identify any predicted LEF/TCF sites (Brannon et al., 1997), suggesting that any Wnt inputs into this particular *cis*-regulatory module are likely to be indirect. Other sites of interest included a GATA site, an LMX site and a Myb site that were all in very close proximity within the putative silencer region. An LMX ortholog (KH.C9.485) is known to be expressed in the *Ciona* notochord (José-Edwards et al., 2011), and functionally important Myb sites have been identified in several *Ciona* notochord CRMs (José-Edwards et al., 2015). GATA TFs have not been implicated in notochord-specific gene expression, but immunostaining suggests there may be *GATA.a* expression in the posterior notochord (Oda-Ishii et al., 2016).

We individually mutated the core motif for each of these TFBSs in the –322 to +13 construct. The GATA and Myb core sequences are almost overlapping, and the LMX site is also nearby, so we took care to mutate these core sequences without affecting the adjacent core sites (Fig. 5A). We cannot exclude, however, that these mutations may have promiscuous effects via important but uncharacterized flanking sequences. Our hypothesis was that one or more of these mutations would have a phenotype similar to the –322 to –165 sequence in which expression in primary notochord is derepressed. We instead found that all 4 mutations led to decreased expression overall, particularly in secondary notochord cells (Fig. 5B). This was particularly evident for the ETS mutation, which is not unexpected given its location outside of the silencer region, the U0126 phenotype and the expression of FGF8/17/18 and FGF9/16/20 in the tip of the tail. It remains unclear, however, whether repression in the primary notochord involves cryptic combinatorial effects of LMX, Myb and/or GATA, or whether there are essential binding sites for unknown factors that have yet to be identified in the –164 to –127 region.

Discussion and Conclusions

Here we characterized the expression patterns and dependencies on Wnt and FGF signaling for three newly identified genes with strongly enriched expression in the posterior notochord. These fell into two classes: *C11.331* and *C12.115* were consistently differentially expressed between primary and secondary notochord, whereas *C8.891* showed a graded expression pattern at several stages extending past the primary/secondary boundary; *C11.331* and *C12.115* were strongly and consistently affected by manipulations of both Wnt and FGF signaling, whereas *C8.891* was strongly affected by ectopic Wnt activation but only modestly affected by FGF inhibition. This indicates that there may be at least two

functionally distinct mechanisms of posterior-specific notochord gene regulation involving multiple signaling molecules expressed in the posterior tip of the tail.

Both *C11.331* and *C12.115* showed a surprisingly complex expression pattern when the canonical Wnt pathway was ectopically activated by BIO treatment. In both cases, the ectopic expression induced by BIO treatment is highly specific to notochord and is not seen in other tissues, confirming that there must be distinct tissue-specific inputs into the expression of these genes. Both expression patterns expanded throughout the notochord upon BIO treatment, but with distinct enrichment in most embryos at both the anterior and posterior tips. This is reminiscent of the wildtype expression pattern for *BCamL* which is similarly enriched in the anterior and posterior notochord. FGFs are expressed in the trunk as well as the posterior tail tip (Hudson and Yasuo, n.d.; Imai et al., 2004), suggesting that they might play a key role in generating these bipolar expression patterns. This is supported by the weak uniform notochord expression of *C11.331* and *C12.115* seen in embryos treated with both BIO and U0126, which shows that induction of tip specific expression by ectopic Wnt pathway activation requires functional FGF signaling. We predict that the combination of both Wnt and FGF signaling at the tail tip is essential for the robust expression of *C11.331* and *C12.115* in posterior notochord, but that *BCamL* likely responds to FGF alone. If this hypothesis is correct, *BCamL* expression should be reduced or eliminated by U0126 treatment but not affected by BIO.

We have also extensively probed the *cis*-regulatory architecture underlying the posterior-specific expression of *C11.331* and find evidence for extensively distributed enhancer function. There are multiple non-overlapping or minimally overlapping regions in the *C11.331* upstream intergenic region that confer posterior notochord-enriched expression in reporter assays. Relatively few ‘shadow/redundant/distributed’ enhancers have been identified to date in *Ciona* (Farley et al., 2016), but they are extremely common in other model systems (Barolo, 2012; Cannavò et al., 2016; Perry et al., 2011) and easily missed in studies focused on rapidly identifying minimal CRM regions. A recent study using ATAC-seq to identify candidate enhancer regions genome-wide suggests that they may be more common in *Ciona* as well (Madgwick et al., 2018).

The functional significance of shadow/redundant/distributed enhancers remains unclear, but multiple non-exclusive hypotheses involving the strength, robustness, tissue-specificity and evolvability of gene expression have been proposed (Barolo, 2012). While not extensively investigated here, our data are consistent with distributed enhancer function being important for both the strength and the tissue-specificity of expression. The smaller constructs that recapitulated the secondary enriched expression seen with the full length –1565 to +13 region tended to have greater ectopic expression and/or weaker expression overall (Fig. 3B).

While the *C11.331* upstream regions showed extensively redundant *cis*-regulatory activity, further dissections of the minimal –488 to +13 regulatory module identified discrete regions with predominant roles in positive and negative regulation. These include a region between –165 and –127 essential for repressing expression in primary notochord, and a region between –245 and –165 inferred to contain essential elements required for both notochord-specific and ectopic expression. The –165 to –127 silencer region is of particular interest,

because it implies that there is a key role for negatively-acting factors that prevent *C11.331* expression in primary notochord. We accordingly searched the –165 to –127 region for TFBSs of interest, but mutations in several predicted sites led to decreased expression in secondary notochord as opposed to increased expression in primary notochord. It remains unclear whether there are yet to be identified sites with clear repressive function, or whether some more complex regulatory scheme might be at work.

It was not clear from the initial *in situ* hybridization experiments whether *C11.331* and *C12.115* are truly specific to secondary notochord, or whether they are induced in the posterior notochord by FGFs and Wnts expressed in the posterior tail tip and coincidentally have an anterior limit of detectable expression that is closely aligned with the primary/secondary boundary. Both the chromogenic and fluorescent *in situ*s made it clear that there is a graded quality to *C11.331* and *C12.115* expression patterns, which are consistently strongest in the posteriormost notochord cells and drop off rapidly towards the primary/secondary boundary. It was unclear, however, whether there was a sharp transition in expression levels at the primary/secondary boundary given the limited sensitivity and modest dynamic range of traditional (non-single molecule) *in situ* hybridization. Quantitative analysis of *C11.331* reporter constructs revealed, however, that although there is graded expression within the secondary notochord, there is a sharp and abrupt transition between very low expression levels in the primary notochord and much higher levels in secondary notochord (Fig. 4A). This suggests that there are inputs into *C11.331* expression based on primary vs secondary notochord fate and that it is not solely based on proximity to Wnt and FGF signals in the tip of the tail.

Fig. 6 shows a provisional model for *C11.331* expression in the secondary notochord. FGF and Wnt signals from a signaling center at the tip of the tail help to induce regionalized *C11.331* expression, which is also dependent on notochord-specific transcriptional regulators (possibly including LMX1 and potential Myb family members) and an as-yet unidentified transcriptional repressor that is predicted to be differentially expressed in primary but not secondary notochord. The identification of this putative primary notochord repressor is of considerable interest, but it will also be important to determine if similar mechanisms are at work in other *C11.331* CRMs as well as in the regulatory regions for other regionally expressed notochord genes. The 1kb upstream regions of *C12.115* and *C8.891* both contain predicted Ets, LMX, GATA and Myb sites that define candidate regions of interest.

Methods:

***Ciona* husbandry and embryology**

Ciona robusta (formerly known as *Ciona intestinalis* type A) (Pennati et al., 2015) were collected in San Diego and shipped to KSU by Marine Research and Educational Products Inc. (M-REP, San Diego, CA). Adult *Ciona* were maintained in a recirculating aquarium. Standard fertilization, dechoriation and electroporation protocols were used (Veeman et al., 2011). Staging is based upon the series of Hotta (Hotta et al., 2007).

Drug treatments: BIO (GSK inhibitor IX, CAS 667463-62-9, Sigma-Aldrich) was dissolved in DMSO at 5 mM and used at a final concentration of 5 micromolar. MEK1/MEK2 inhibitor U0126 (CAS109511-58, Sigma-Aldrich) was dissolved in DMSO at 4 mM and used at a final concentration of 4 micromolar. DMSO alone (at 1:1000 dilution) was used as a control. Drugs were added to artificial seawater (ASW) at Hotta stage 12 and embryos were grown in drug-treated ASW until fixation at Hotta stage 23.

***In situ* hybridization**

Probe synthesis, embryo collection, *in situ* hybridization and imaging were performed essentially as described previously (Reeves 2014), except that embryos for fluorescent *in situ* were electroporated with 60 micrograms *Bra*-hCD4:mCherry (Gline et al., 2015) to visualize notochord cell boundaries post-hybridization. mCherry antibody (Biovision, #5993) was included at 1:500 during the anti-DIG-AP incubation. After SIGMA-FAST FastRed staining (Sigma-Aldrich; F4648) was stopped by washing 5 times in PBS-Tween20, embryos were incubated overnight at 4°C with 1:1000 anti-rabbit AlexaFluor 488 (Invitrogen, A-11034). Embryos were washed 5 times in PBS-Tween20, then mounted, cleared, imaged and fluorescence was quantified in cells 30–40 as described previously (Reeves et al., 2014). All fluorescence intensities were background subtracted and then normalized to a 0–1 scale. Expression data was centered on the primary/secondary boundary and scaled to a common length.

The *Brachyury in situ* probe was previously described (Reeves et al., 2014). Regions of *C11.331*, *C12.115* and *C8.891* were amplified from cDNA and cloned into pBSII-SK(-), using the following primers: *C11.331* (993bp, Forward: ACGGGACTCACACAACCTCC, Reverse: GTCTCCAATCGCTTGCTGTT); *C12.115* (865bp, Forward: ACGCCATTAACACCGGTTTC, Reverse: CAAATGTTTGTAGAAAAGTATTGAC), *C8.891* (900bp, Forward: GTGCTGATGCCAAGAATGC, Reverse: GTTTCACACAGCTGGTAGGC).

Reporter cloning and mutagenesis

The genomic region spanning from -1565 to +13 of KH2012:KH.C11.331.v1.A.SL2-1 was PCR amplified from genomic DNA (Forward primer: TAAAATGGCGCGCCAGGTGCCACAAATAAACC; Reverse primer: CCTCCGTCTAGACCTATTTGTCCTTCTGAAATAACAG and cloned into the *AscI* and *XbaI* sites of pX2+bpFOG>UNC76:Venus (Stolfi et al., 2015). All subsequent reporters were subcloned from this original plasmid (see Table S5 for primers and genomic coordinates). TFBS single mutants were generated through the Q5 Site-Directed Mutagenesis Kit (E0554S, NEB), following manufacturer directions. Mutagenesis primers are listed in Table S5. The *Bra*-H2B:HA reporter was generated by standard Gateway cloning of pENTR-H2B into pSP72BSSPE-SwaI::RFA-HA (Roure et al., 2007) A 2.2kb *Brachyury* enhancer was cloned into the plasmid's *XhoI*/*HindIII* sites.

Reporter expression, staining, and imaging

Fertilized dechorionated eggs were co-electroporated with 30 µg *Bra* > H2B:HA plasmid and 60 µg *C11.331* reporter plasmid (region of interest-bpFOG>Venus). At least three

separate replicates were performed for each reporter. Embryos were fixed at Hotta stage 21–22, stained and prepared for imaging as previously described (Carlson et al., 2015). GFP polyclonal antibody (Invitrogen, A-11122) was used at 1:000 and HA-Tag mouse monoclonal antibody (Cell Signaling Technology, 2367S) at 1:750. Secondary antibodies (Invitrogen; anti-mouse AlexaFluor 488, A-11029 and anti-rabbit AlexaFluor 555, A-21420) were used at 1:1000, and AlexaFluor 633 Phalloidin (Invitrogen, A22284) was used at 1:150. Embryo identities were blinded prior to mounting and clearing embryos. Mounted embryos were imaged on a Zeiss 700 laser scanning confocal microscope, using a 40× 1.3NA objective. Z-stacks of each embryo were collected at a slice interval of 0.5 μm and a pixel size of 0.24 μm, using consistent settings for laser power, PMT gain and scan speed. Embryos were selected for imaging based on expression of the *Bra*>H2B:HA control in both primary and secondary notochord, but without respect for the expression of the *C11.331* reporter plasmid.

Quantitative reporter analysis

Confocal stacks of embryos electroporated with the indicated reporter plasmids were analyzed interactively using FIJI/ImageJ (Schindelin et al., 2012). The tail is rarely flat enough or sufficiently parallel to the coverslip to visualize the entire notochord in one 2D image. To allow a simple 2D analysis of reporter intensity, the notochord was first computationally flattened in the Z axis to bring all the *Bra*>H2B:HA labeled nuclei into a single plane. This involved reslicing the image to view the image volume as XZ or YZ slices, manually tracing a polyline that followed the notochord in Z, and then reslicing again along the polyline to reconstruct a flattened plane through the full length of the notochord. Embryos that could not be cleanly resliced in this manner to generate a clear image of the embryo were rejected from quantitative analysis. A 40 pixel wide polyline was then traced along the notochord's curvature in X and Y to capture the full width of each labeled nucleus from cell 1 to cell 40. Intensity data from the Venus and HA reporter channels was collected along that line using the 'plot profile' function. All of these steps used standard FIJI/ImageJ tools.

We then derived a normalized expression metric implemented as a MATLAB function. Inputs to the function include the AP intensity profiles of Venus and HA, manually estimated values for Venus and HA background levels, and the inferred positions of the primary/secondary notochord boundary and the cell 36/37 boundary (anterior versus posterior secondary notochord). The intensity profiles were segmented into the three regions of interest. For each region, a normalized expression score was generated by dividing background-subtracted reporter>Venus intensity by control background-subtracted *Bra*>HA intensity, but only for the positions with greater than threshold levels of *Bra*>HA expression.

Wilcoxon signed-rank tests were used to compare expression levels between notochord regions for the quantitative analysis. ANOVA followed by Tukey's Honest Significant Difference post-hoc test was used to compare differences in expression between different reporter constructs for each notochord subregion to better control for multiple comparisons. All statistical tests used standard MATLAB functions. We used the Python Pandas, Numpy, Matplotlib and Seaborn packages for plotting and data visualization.

Qualitative reporter analysis

Confocal stacks of embryos from the reporter dissections were scored for expression in the primary notochord, anterior four cells of the secondary notochord, and posterior four cells of the secondary notochord on a scale from zero (no visible expression) to four (oversaturated expression). Scores were assigned based on the cell in the notochord region with the strongest Venus expression that also expressed *Bra*>H2B:HA. Scores were also given for ectopic expression in CNS, epidermis, endoderm, muscle, and mesenchyme, but without the requirement for *Bra*>H2B:HA expression in the same cell. Little evidence was seen for distinct tissue-specific silencers outside the notochord, so a combined ectopic expression score was derived by averaging the scores across the 5 tissues examined. Scoring was performed separately and without knowledge of the specific secondary reporter plasmid by both MH and CB prior to reconciling any score differences and recording consensus scores for each embryo. For simpler presentation, we averaged the anterior and posterior secondary notochord scores into a single combined value.

TFBS Analysis

The predicted ETS, GATA and Myb sites were identified using TRANSFAC PWMs (Matys et al., 2006) via the LASAGNA web tool (http://bioGRID-lasagna.engr.uconn.edu/lasagna_search/) (Lee and Huang, 2013). The LMX site was identified by manually searching the minimal region of interest for matches to the core sequence identified by SELEX against *Ciona* Lmx1A (Brozovic et al., 2018). We also searched for matches to SELEX-derived motifs for Brachyury (shared by Tbx2/3) and FoxAa but did not locate any within this interval.

Resources

Information on key resources are provided in the KRT table.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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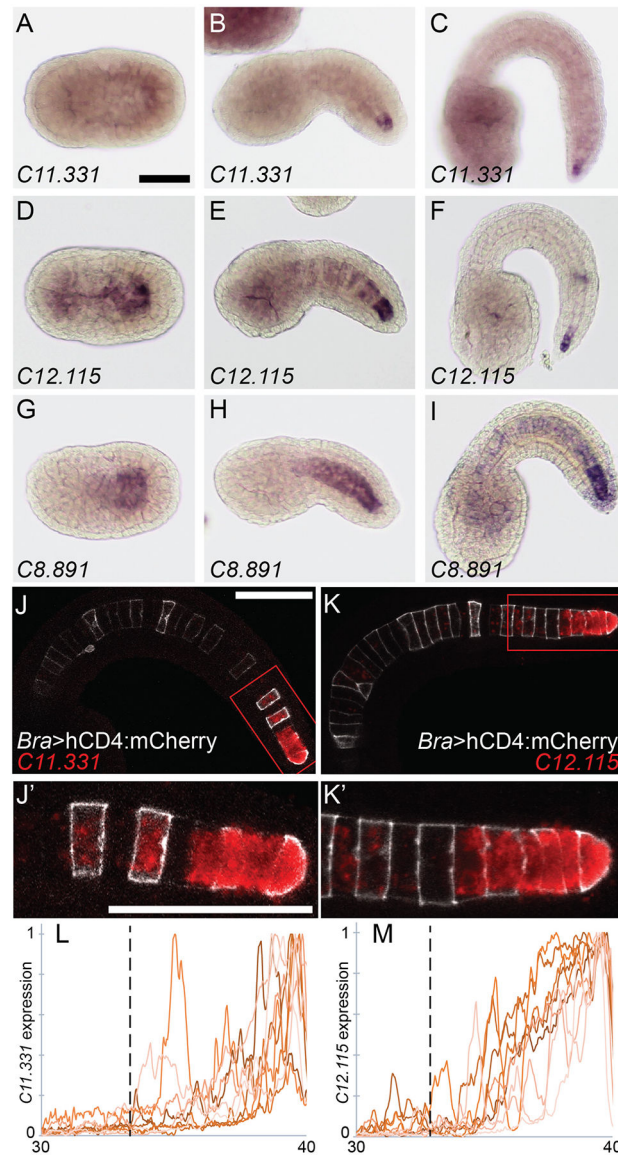


Fig. 1. Secondary-enriched notochord genes.

In situ hybridization of probes against (A-C) *KH.C11.331*, (D-F) *KH.C12.115* and (G-I) *KH.C8.891*. Expression was tested during early intercalation at Hotta stage 16 (A, D, G), late in intercalation at Hotta stage 20 (B, E, H) and during notochord elongation at Hotta stage 23 (C, F, I). J-K) Fluorescent *in situ* hybridization against *C11.331* (J) and *C12.115* (K) in red at Hotta stage 22. Embryos were electroporated with *Bra>hCD4:mCherry* (white). J'-K') Closeup of secondary notochord region indicated by red box in (J,K). L-M) Background subtracted and normalized expression of *C11.331* (L) and *C12.115* (M) in notochord cells 30–40. Nine embryos for each gene are shown. The location of the primary/secondary border is indicated by the vertical dotted line. Scale bars = 50 microns.

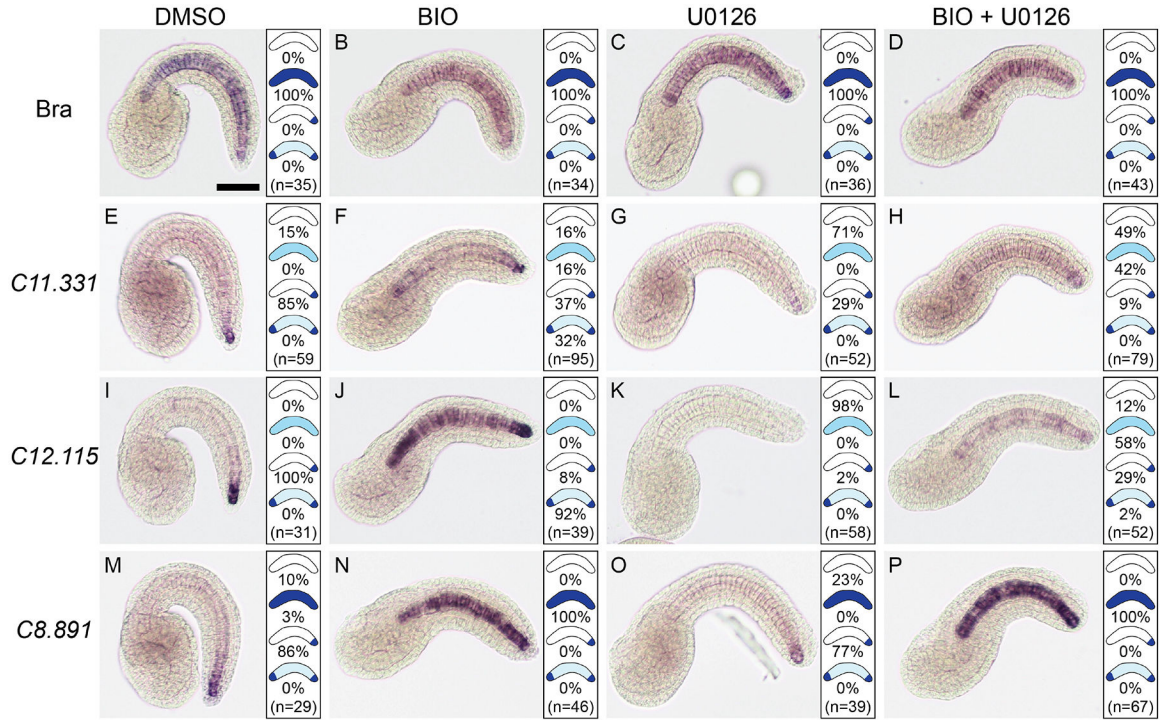


Fig. 2. Wnt and FGF signaling regulate posterior-enriched notochord expression. *In situ* hybridization against *Brachyury* (A-D), *C11.331* (E-H), *C12.115* (I-L) and *C8.891* (M-P) on Hotta stage 23 embryos treated starting at mid gastrula (Hotta stage 12) with 1:1000 DMSO (A, E, I, M), 5 μ M Wnt activator BIO (B, F, J, N), 4 μ M FGF inhibitor U0126 (C, G, K, O) or BIO + U0126 (D, H, L, P). Representative images of the most common expression pattern for each gene/treatment are shown. Panels to the right of each image show the distribution of expression patterns observed. Scale bar = 50 microns.

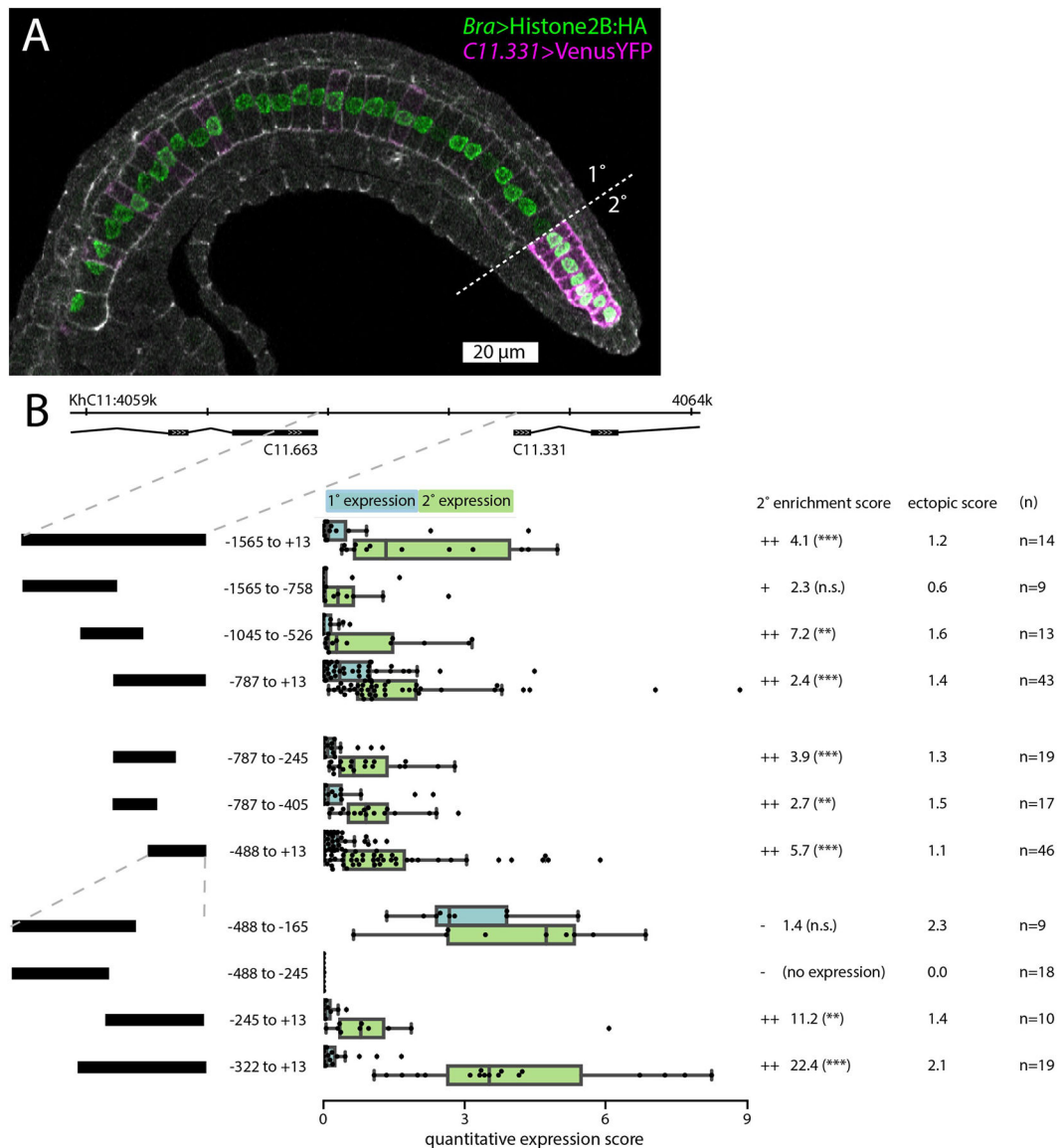


Fig. 3. Cis-regulatory analysis of the *C11.331* upstream intergenic region.

A: Representative embryo expressing *C11.331* (-787 to +13)>Venus and *Bra*>Histone2B:HA fixed during tail elongation at Hotta stage 22. Cell cortices are labeled with Phalloidin-Alexa633 (white). Primary/secondary notochord boundary indicated with dashed line. Anterior is to the left and dorsal is to the top of the image. **B:** Summary statistics from reporter assays for the full-length *C11.331* upstream region and a series of truncation mutants. Construct coordinates refer to the predicted transcriptional start site for *C11.331*. *C11.331* quantitative expression score distributions are shown for primary (blue) and secondary (green) notochord. The superimposed box plots for each distribution show the median flanked by the interquartile range, with whiskers extending to the full range excluding outliers >1.5X IQR. Expression differences between notochord regions for each reporter are shown as enrichment scores, calculated as the mean secondary expression divided by the mean primary expression. ++ indicates that the construct was

enriched in secondary notochord with statistical significance <0.05 . + indicates that the construct showed signs of enrichment in secondary notochord but it was not statistically significant. – indicates that the construct was not enriched in secondary notochord. Secondary enrichment was tested by one-tailed Wilcoxon signed-rank tests (*, $p<0.05$; **, $p<0.005$; ***, $p<0.0005$). The ectopic expression scores represent the average expression in 5 non-notochord tissues using a 0–4 qualitative expression scale described in the methods. For a tissue-specific breakdown of the scores, see Tables S3,S4.

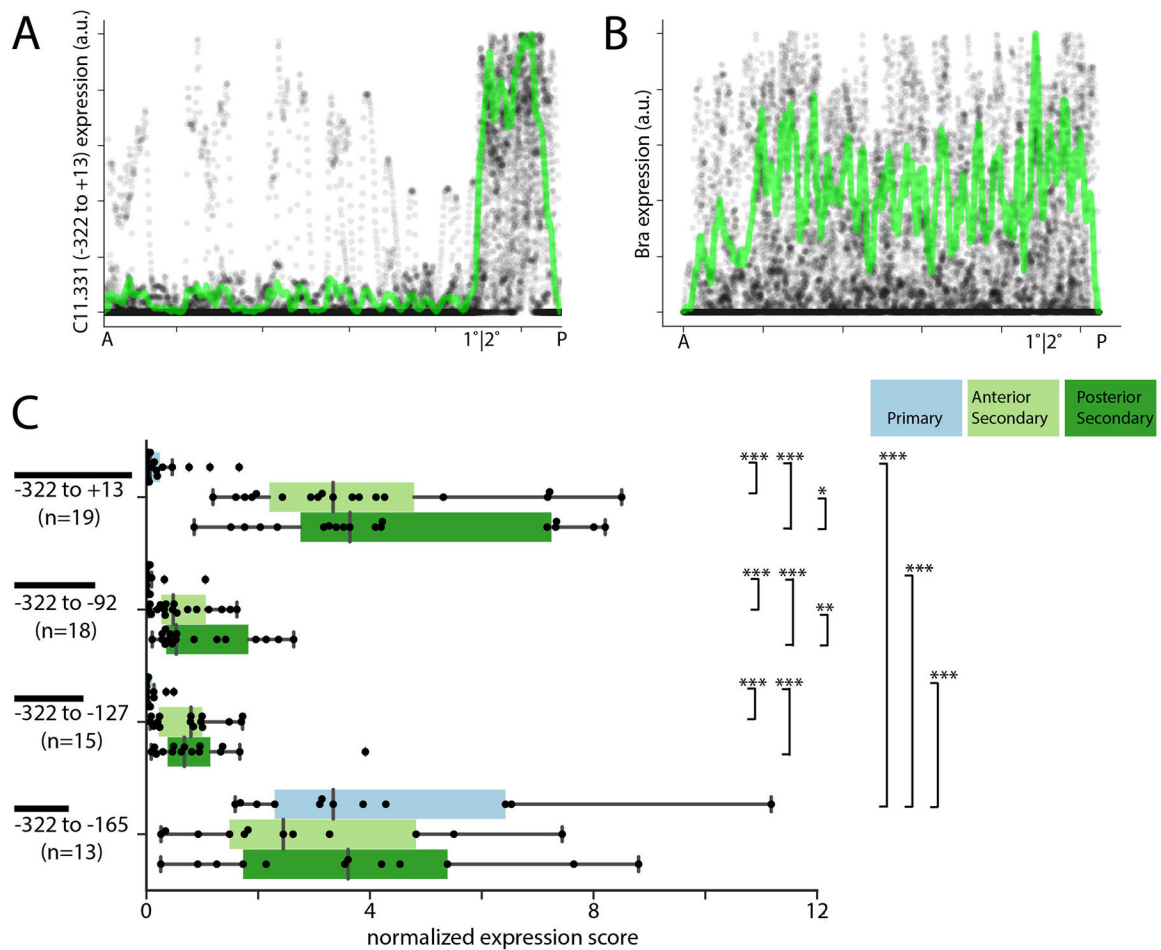


Fig. 4. Quantitative analysis of the *C11.331*(-322 to +13) CRM.

A-B) Quantitative traces of *C11.331*(-322 to +13) (A) and *Bra* (B) expression from 19 embryos as a function of anterior-posterior position. Expression data from each embryo was scaled in intensity to {0 1}, scaled to a common length, aligned by the primary/secondary notochord boundary and plotted as transparent gray to help visualize overlapping data points. Normalized mean intensity is overlaid in green. C) *C11.331* quantitative expression score distributions are shown for primary (blue) anterior secondary (light green) and posterior secondary (dark green) notochord. The superimposed box plots for each distribution show the median flanked by the interquartile range, with whiskers extending to the full range excluding outliers >1.5X IQR. Expression differences between notochord regions for a given reporter were tested by one-tailed Wilcoxon signed-rank tests. Expression differences for each notochord region between the different reporter constructs were tested by ANOVA followed by Tukey's Honest Significant Difference post-hoc test (*, $p < 0.05$; **, $p < 0.005$; ***, $p < 0.0005$).

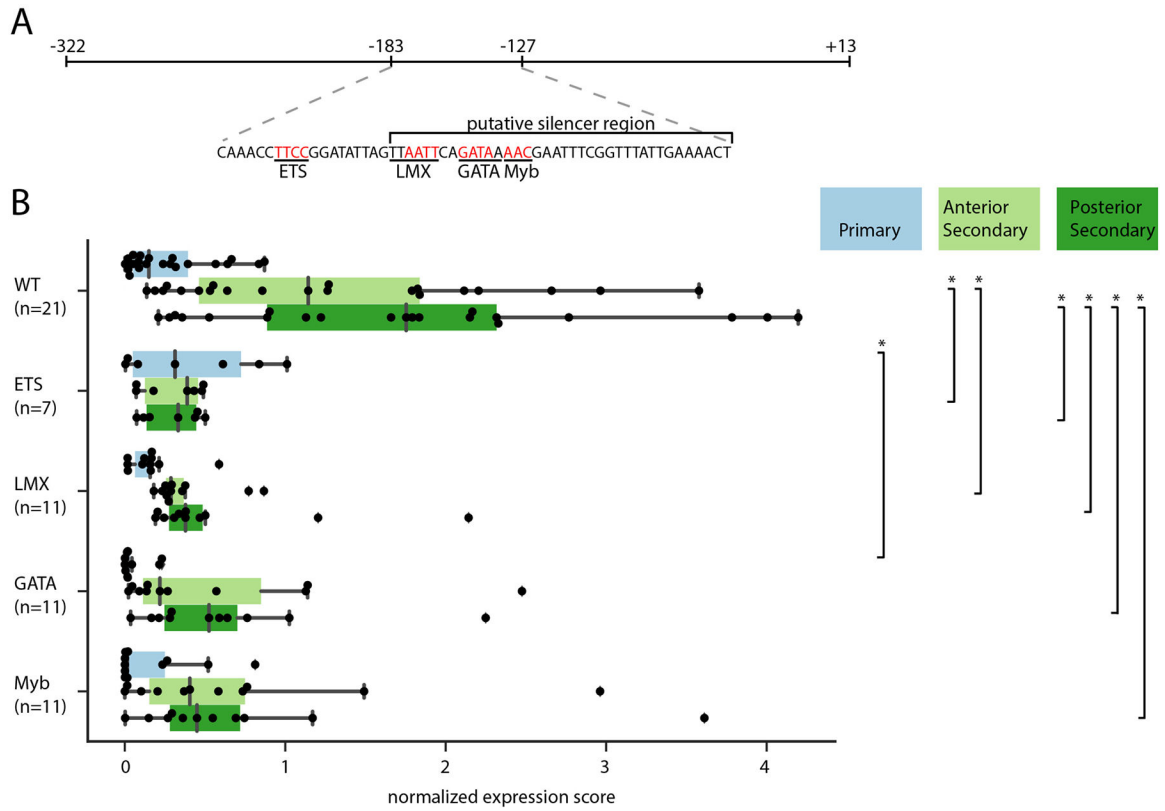


Fig. 5. Mapping TFBSs in the *C11.331* (-322 to +13) CRM.

A) Schematic of *C11.331* (-322 to +13) with the sequence of the -183 to -127 region shown and the -165 to -127 putative silencer region indicated. Core binding sites for the transcription factors ETS, LMX, GATA, and Myb are underlined, with the specific bases mutated in derivative constructs shown in red. B) Quantitative analysis of regionalized notochord expression for the *C11.331*(-322 to +13) construct and derivatives in which each of these predicted TFBSs have been mutated. Quantitative expression scores for individual embryos are shown for primary notochord (blue) anterior secondary (light green) and posterior secondary (dark green). The superimposed box plots for each distribution show the median flanked by the interquartile range, with whiskers extending to the full range excluding outliers >1.5X IQR. Between-reporter differences in expression for each notochord region were tested by ANOVA followed by Tukey’s Honest Significant Difference post-hoc test (*, $p < 0.05$).

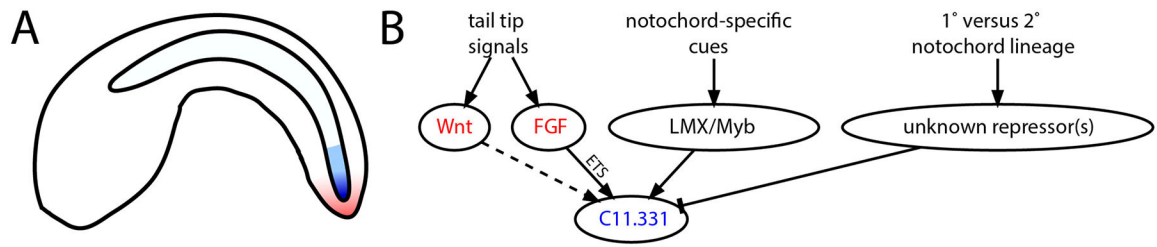


Fig. 6. Regulatory network model for *C11.331* expression.

A) Schematic of a *Ciona* embryo showing Wnt and FGF expression in the posterior tail tip (red) and *C11.331* expression in blue. *C11.331* expression is both highly enriched in secondary notochord compared to primary notochord and also graded within the secondary notochord. B) Simple network model of inferred inputs into *C11.331* expression.

KEY RESOURCES TABLE

Reagent or resource	Source	Identifier
Antibodies		
Rabbit polyclonal anti-mCherry	Biovision	Cat#5993
Goat anti-rabbit polyclonal AlexaFluor488	Invitrogen	Cat#A11034
Goat anti-mouse polyclonal AlexaFluor488	Invitrogen	Cat#A11029
Goat anti-rabbit polyclonal AlexaFluor555	Invitrogen	Cat#A22420
Rabbit polyclonal anti-GFP	Invitrogen	Cat#A11122
Mouse monoclonal anti-HA	Cell Signaling Technology	Cat#2367S
Bacterial and Virus Strains		
n/a		
Biological Samples		
n/a		
Chemicals, Peptides, and Recombinant Proteins		
GSK inhibitor IX (BIO)	Sigma-Aldrich	Cat#361550; CAS667463-62-9
U1026	Sigma-Aldrich	Cat#662005; CAS109511-58-2
SIGMA-FAST FastRed	Sigma-Aldrich	Cat#F4648
AlexaFluor633 Phalloidin	Invitrogen	Cat#A22284
Critical Commercial Assays		
n/a		
Deposited Data		
<i>Ciona</i> SELEX data	ANISEED	https://www.aniseed.cnrs.fr/
Experimental Models: Cell Lines		
n/a		
Experimental Models: Organisms/Strains		
Wild <i>Ciona robustus</i>	M-REP, San Diego, CA	N/A
Oligonucleotides		
See Table S1		
Recombinant DNA		
<i>Bra</i> -hCD4:mCherry	Gline et al, 2015	N/A
All <i>C11.33I</i> >UNC76:Venus reporter plasmids	This paper	N/A
<i>Bra</i> -H2B:HA	This paper	N/A
Software and Algorithms		
MATLAB	The Mathworks, Inc	Release 2012b
FIJI/ImageJ	Schindelin et al, 2012	https://fiji.sc/
LASAGNA-GRID	Lee & Huang, 2013	http://biogrid-lasagna.engr.uconn.edu/lasagna_search

Reagent or resource	Source	Identifier
Other		

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