

A missing link in the sea urchin embryo gene regulatory network: *hesC* and the double-negative specification of micromeres

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Specification of sea urchin embryo micromeres occurs early in cleavage, with the establishment of a well defined regulatory state. The architecture of the gene regulatory network controlling the specification process indicates that transcription of the initial tier of control genes depends on a double-negative gate. A gene encoding a transcriptional repressor, *pmar1*, is activated specifically in micromeres, where it represses transcription of a second repressor that is otherwise active globally. Thus, the micromere-specific control genes, which are the target of the second repressor, are expressed exclusively in this lineage. The double-negative specification gate was logically required from the results of numerous prior experiments, but the identity of the gene encoding the second repressor remained elusive. Here we show that *hesC* is this gene, and we demonstrate experimentally all of its predicted functions, including global repression of micromere-specific regulatory genes. As logically required, blockade of *hesC* mRNA translation and global overexpression of *pmar1* mRNA have the same effect, which is to cause all of the cells of the embryo to express micromere-specific genes.

skeletogenic micromeres | transcriptional repression

The genomic regulatory code for specification of endomesoderm in the sea urchin embryo is represented as a gene regulatory network (GRN), which explains the mechanism by which distinct regulatory states are deployed in different territories of the developing embryo (for reviews, see refs. 1–4; for current version see <http://supg.caltech.edu/endomes>). One portion of this GRN pertains to the specification of the micromeres, which arise at the unequal fourth cleavage at which the four micromeres are segregated off from the vegetal pole of the egg. The large daughter cells of the micromeres arising at the next cleavage are the founder cells of the skeletogenic micromere lineage. This lineage is the sole normal source of the embryonic biomineral skeleton, a distinct synapomorphic feature of echinoid embryos and larvae, and it also produces essential short-range signals required for other aspects of endomesoderm specification (5–7). Three particular developmental events that are relevant for what follows are (i) the expression of the Delta-signaling ligand on the surfaces of the micromere descendants during the early blastula stage (Fig. 1A); (ii) their ingression into the blastocoel at the late blastula stage (Fig. 1B), after which they are known as primary mesenchyme cells; and (iii) their expression of the biomineralization and cytoskeletal genes, which enable them to generate the skeleton (Fig. 1C) (8).

Immediately after the fourth-cleavage micromeres are born, they express a gene, *pmar1*, in response to maternally localized factors (9). This gene encodes a transcriptional repressor of the paired homeodomain family. In the GRN model, *pmar1* serves as the linchpin of a proposed double-negative gate controlling the institution of the micromere regulatory state. The second component of this gate is predicted to be a *pmar1* target gene that encodes another transcriptional repressor. This gene must also be zygotically expressed, but it would be transcribed everywhere in the embryo except in the micromere lineage, where it

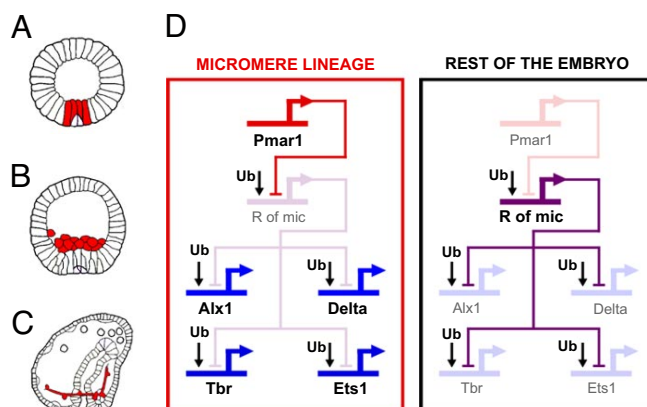


Fig. 1. Key elements of the GRN model for the early specification of the skeletogenic micromere lineage. The model is based on ref. 9, with subsequent updates (10), as reviewed in refs. 1 and 4. (A–C) Sea urchin embryo drawings (adapted from ref. 2) at the early blastula stage (12–15 h after fertilization; A), at the mesenchyme blastula stage (24 h; B), and at the late gastrula stage (48 h; C). The cells of the skeletogenic micromere lineage at each stage are depicted in red. (D) The GRN model (corresponding to cleavage and blastula stages). Active genes are represented in strong color and bold font. Inactive genes are represented in dim color. Within the micromere lineage, *pmar1* is active and represses the predicted gene *r of mic*. The *delta*, *alx1*, *ets1*, and *tbr* genes are allowed to be zygotically expressed in this domain. In the rest of the embryo, *r of mic* keeps *delta*, *alx1*, *ets1*, and *tbr* silent.

is subject to repression by *Pmar1*. There are eight targets predicted for it in the GRN, of which the most important for present purposes are the genes encoding the Delta ligand and three regulatory genes, *tbr*, *ets1*, and *alx1*. These three genes lie upstream of all the rest of the micromere regulatory apparatus. In this manner, the double-negative gate would ensure expression of this apparatus exclusively in the micromere lineage. Because its identity was unknown, the second repressor has been referred to in the GRN as Repressor of Micromeres (R of mic). Its existence and properties are specifically implied by the two following perturbation experiments (9, 10). First, if expression of *pmar1* is forced to occur globally (by injection into the egg of the mRNA), then the *delta*, *tbr*, *ets1*, *alx1*, and downstream genes are transcribed in all cells of the embryo, and all cells thereby adopt

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Abbreviations: DIG, digoxigenin; DNP, dinitrophenol; GRN, gene regulatory network; MASO, morpholino antisense oligonucleotide; MOE, mRNA overexpression; WMISH, whole-mount *in situ* hybridization.

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skeletogenic micromere lineage fate. Second, exactly the same outcome follows if an mRNA encoding a dominantly repressive Engrailed fusion of the Pmar1 protein is injected. It follows that the *pmar1* gene product naturally acts as a repressor (also indicated by its sequence); that *delta*, *tbr*, *ets1*, and *alx1* are controlled by ubiquitous activators; and that localization of expression of these genes to the micromere lineage in normal embryos depends on their repression by R of mic everywhere else in the embryo (Fig. 1D).

To prove the existence of the double-negative gate for micromere lineage specification in the GRN model, it is necessary to find the gene playing the role of the predicted R of mic and establish that its expression and functions are also as predicted. The *r of mic* gene should encode a transcriptional repressor, and it should have three distinct characteristics: (i) Its zygotic expression should be transcriptionally repressed by Pmar1; (ii) it should be zygotically expressed everywhere except in the micromere lineage by the time zygotic expression of *delta*, *alx1*, *ets1*, and *tbr* starts; and (iii) the outcome of knocking down its expression should be similar to forcing global Pmar1 expression (i.e., all cells of the embryo should adopt micromere lineage specification and express *delta*, *alx1*, *ets1*, and *tbr*).

Results

Genomic Screen for Candidate *r of mic* Genes. The *Strongylocentrotus purpuratus* genome sequence enabled consideration of all sea urchin genes encoding transcription factors in our search for *r of mic*. The total number of annotated transcription factors in this genome, excluding C₂H₂ zinc finger genes, is 283 (11), and the total number of predicted C₂H₂ zinc fingers (some of which encode transcription factors) is 377 (12). The levels of mRNA expression at several developmental time points were measured for all of these 660 genes (12–16). We selected as *r of mic* candidates all putative regulatory genes for which at least 200 transcripts were detected per embryo at 12 h after fertilization, when *delta*, *alx1*, *ets1*, and *tbr* are all zygotically transcribed. This threshold is conservative (low) given that *r of mic* must be expressed in most of the embryo, or in 100 to 150 cells, at this time. This selection resulted in a list of ≈100 candidate genes. We excluded those that are maternally and not zygotically expressed up to 12 h after fertilization and all previously studied transcription factors for which enough information was available to confirm that they could not be *r of mic*. The surviving list now contained 46 candidates [see supporting information (SI) Table 1].

Because *r of mic* should be transcriptionally repressed by Pmar1, we screened the 46 candidate genes for down-regulation on forced expression of Pmar1 in the whole embryo [Fig. 2 and SI Fig. 6; mRNA overexpression (MOE)]. The effect of *pmar1* MOE on the level of transcripts of each *r of mic* candidate gene was measured at 9 and 12 h after fertilization by using quantitative PCR. The *delta* gene was included in the screen as a control. As expected, in the two experiments performed, *delta* was significantly up-regulated (≥3-fold changes in transcript levels were considered significant) at both 9 and 12 h after fertilization (Fig. 2). This result indicated that *r of mic* must have been down-regulated at both time points in these two experiments. Five of the 46 regulatory genes tested were found to be significantly down-regulated at both time points in the two experiments performed. These genes were *six3*, *smadIP*, *awh*, *hesC*, and *foxJ1* (Fig. 2).

Among these five transcriptional regulatory genes, *hesC* particularly caught our attention. Its level of mRNA expression at 9 and 12 h is highest of all five (data not shown). In addition, HesC is a basic helix–loop–helix transcription factor belonging to the Hairy/E(spl) family, and almost all transcription factors of this family are known to function as repressors (17). That HesC belongs to this family is supported by a phylogenetic analysis (16) and by the fact that it contains the two characteristic domains of

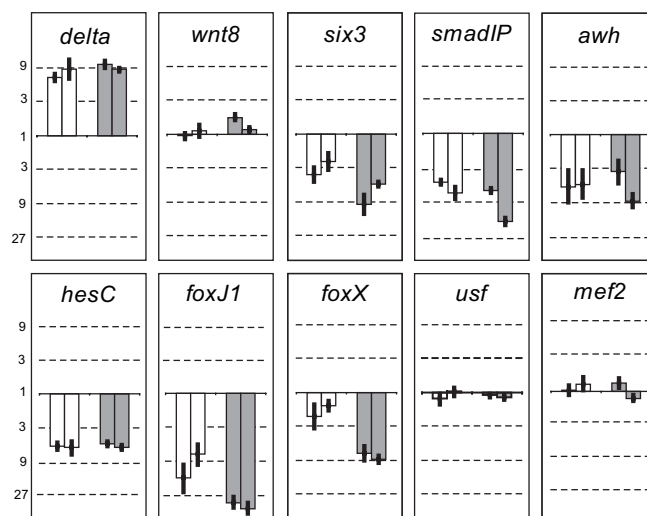


Fig. 2. *pmar1* MOE screen. Graphs showing fold change in mRNA expression for *r of mic* candidate genes on overexpression of *pmar1* mRNA. *delta* and *wnt8* were used as positive and negative controls, respectively. A fold change of 1 (solid line) indicates no change. The numbers situated above 1 indicate a fold increase, whereas the numbers situated below 1 indicate a fold decrease (in logarithmic scale). A 3-fold or greater change was considered to be significant. Open and filled bars represent data from samples at 9 and 12 h of development, respectively. For each type of bar, the two bars correspond to two independent batches of embryos. Error bars represent the standard deviation from three independent measurements on the same sample. Results for 8 of the 46 *r of mic* candidate genes are shown here (see SI Fig. 6 for data on the remaining 38 candidates).

the family: the C-terminus WRPW motif (used to recruit TLE/Grg/Groucho and mediate transcriptional repression) (18, 19) and the Orange domain. We therefore focused on *hesC*.

Temporal and Spatial Expression of *HesC*. The spatial and temporal patterns of expression predicted for the *r of mic* gene are unique. The time course of *hesC* expression was determined at 1- to 2-h intervals by means of quantitative PCR (Fig. 3A). This experiment showed that *hesC* is maternally expressed, but only at low levels. The level of *hesC* transcript then increases steeply between 8 and 12 h after fertilization, indicating zygotic transcription. To compare the temporal expression of *hesC* to that of up- and downstream genes in the double-negative gate, we measured the levels of *pmar1* and *delta* mRNA in the same embryo samples (Fig. 3A). As would be expected for *r of mic*, the zygotic expression of *hesC* starts before *delta* is detected, and it occurs while *pmar1* mRNA is present.

Whole-mount *in situ* hybridization (WMISH) provided strong evidence. At 8 h, the steep zygotic expression of *hesC* had just started and at this time, *hesC* mRNA was found essentially everywhere in the embryo, including the micromere lineage (Fig. 3F; compare to control in Fig. 3B). The two small cells at the vegetal pole of the embryo, which showed weaker staining than the rest of the embryo, are the small micromeres, that do not belong to the skeletogenic micromere lineage, which is the subject of this article. At 12 h, the steep zygotic increase in *hesC* expression attained its plateau value (Fig. 3A), and *delta* mRNA was already present. The expression of *hesC* had changed dramatically, in that this transcript disappeared from a set of 12 cells at the vegetal pole (Fig. 3C and G), whereas it continued to be expressed everywhere else. Exactly 12 cells expressed *delta* mRNA at this time (Fig. 3D and H), and the same number of cells are now in the micromere lineage. To confirm that the 12 cells lacking *hesC* mRNA expression corresponded to the 12 micromere lineage cells, we performed double-WMISH. As

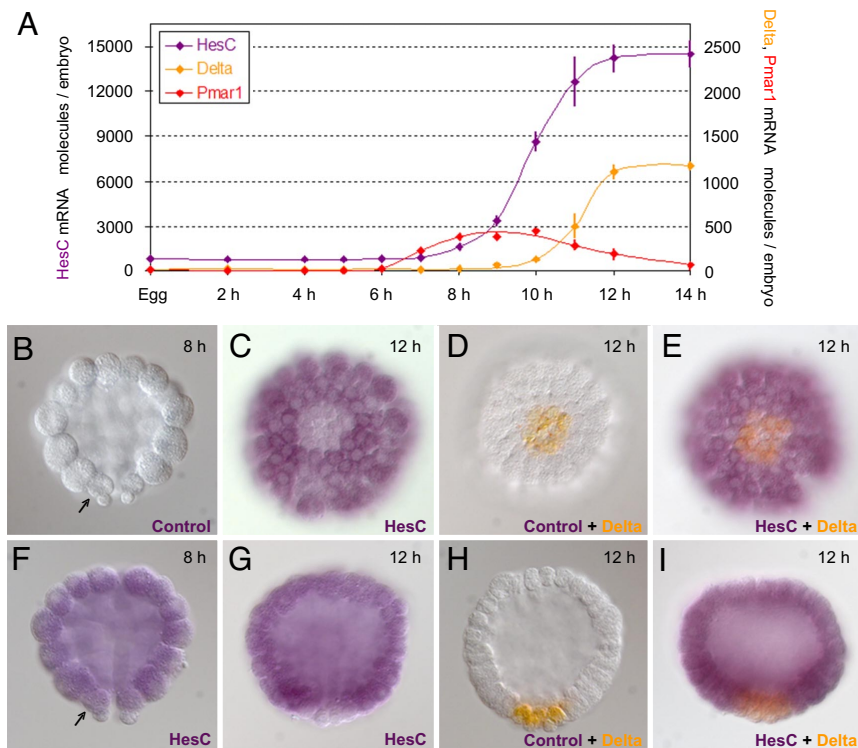


Fig. 3. HesC temporal and spatial expression pattern. (A) Measurements of *hesC* mRNA molecules per embryo (purple) are compared with those of *pmar1* (red) and *delta* (orange) at the indicated developmental time points. Error bars represent the standard deviation from three individual measurements on the same sample. (B–I) Images of embryos on which WMISH (B, C, F, and G) or double-WMISH (D, E, H, and I) was performed. The developmental stage of each embryo is indicated at the upper right corner. (B and F–I) Side views, with a vegetal side at the bottom. (C–E) Vegetal views. The arrows in B and F point to one of the two visible skeletogenic micromere cells. Probes used are indicated at the lower right corner. Control, probe used to control for nonspecific staining.

shown in Fig. 3 E and I, every cell of the embryo expressed either *hesC* (purple) or δ (orange), but no cell expressed both genes. Therefore, zygotic expression of *hesC* had occurred everywhere in the embryo except the micromere lineage, precisely as predicted for *r of mic*.

Functional Analysis of *hesC*. The predicted function of *r of mic* is to repress micromere lineage specification. Thus, if *hesC* is *r of mic*, blocking its translation should result in all cells of the embryo becoming specified similarly to skeletogenic micromeres, the same as when global *pmar1* expression is forced to occur (9). We used a morpholino antisense oligonucleotide (MASO) targeting *hesC* mRNA for this experiment. The striking effect of this perturbation on the morphology of the developing embryos is shown in Fig. 4. Up to the blastula stage, *hesC* MASO embryos were indistinguishable from unperturbed embryos (Fig. 4 A and C). In both, ingress of primary mesenchyme cells into the blastocoel started ≈ 20 h after fertilization (data not shown). However, in unperturbed embryos, primary mesenchyme cell ingress had been completed by 24 h after fertilization (Fig. 4 B), whereas in *hesC* MASO embryos, ingress of cells continued until the blastocoel was essentially full (Fig. 4 D). All, or almost all, cells of *hesC* MASO embryos thus behave in a way normally unique to the micromere lineage. Importantly, at all three stages, *hesC* MASO embryos look strikingly similar to *pmar1* MOE embryos (Fig. 4 C–F; data not shown for 20-h stage).

We next assessed the effect of HesC MASO perturbation on the levels of mRNA of *delta*, *alx1*, *ets1*, and *tbr*. If *hesC* is *r of mic*, the prediction (Fig. 1 D) is that these genes will now be allowed to be expressed in all cells, and their level of transcript should therefore increase, as occurs in *pmar1* MOE embryos (9, 10). Fig.

5 and SI Table 2 show this result. By 12 h after fertilization, the amount of transcript of *delta* and *alx1* had increased 4- to 7-fold above normal in the two experiments performed (Fig. 5 A), and by 24 h, that of *ets1* and *tbr* had similarly increased (SI Table 2). The level of expression of *pmar1* was not affected, indicating that the up-regulation of these genes was not caused by any change in *pmar1* (Fig. 5 A).

The derepression of micromere lineage specification occurs in all cells of *hesC* MASO embryos. This finding is illustrated for the *delta* marker, as shown in Fig. 5 B–E. In unperturbed embryos, *delta* mRNA is localized to the micromere lineage (Fig. 5 B and D), whereas in *hesC* MASO embryos, it is detected throughout the whole embryo (Fig. 5 C and E). Thus, HesC functions to repress micromere lineage specification in all cells other than the micromere lineage, the defining characteristic of the predicted *r of mic*.

Discussion

The GRN Model Prediction and the Evidence. As regions of a GRN approach completion, the levers of logic can be used to generate precise predictions of missing components. As described elsewhere, the portion of the sea urchin endomesoderm GRN that pertains to specification and initial differentiation of the skeletogenic micromere domain now includes every regulatory gene predicted in the genomic sequence that is expressed specifically in the micromere lineage up to the onset of gastrulation (see <http://sugp.caltech.edu/endomes>), although some linkages among these genes may remain to be determined. Further, the GRN does not identify genes expressed ubiquitously in the embryo, which provide inputs to micromere genes, and it includes only a sample of the terminal skeletogenic differentiation genes. Nonetheless, it should include all, or almost all,

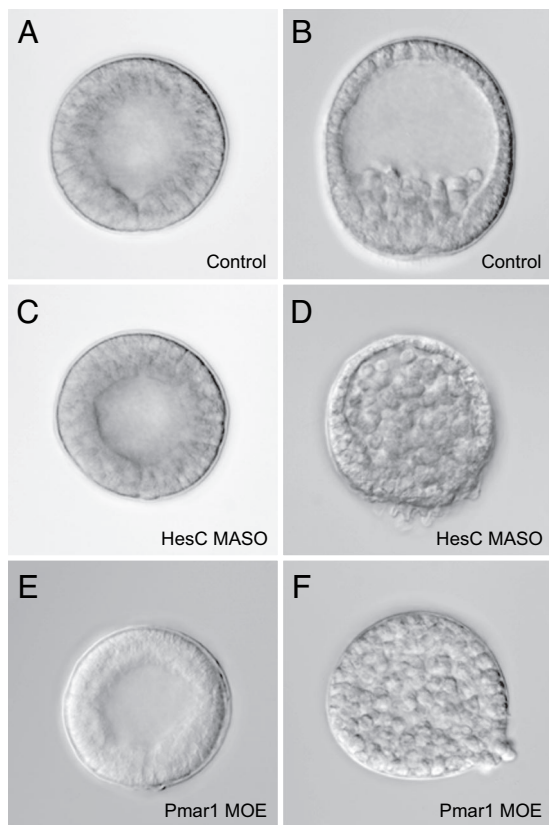


Fig. 4. Morphology of HesC MASO embryos. (A–D) Images of embryos that were either unperturbed (A and B) or perturbed (C and D) by HesC MASO. (E and F) Pmar1 MOE embryos from a different batch are also shown for comparison. (A, C, and E) Blastula stage embryos 16 h after fertilization. (B, D, and F) Late mesenchyme blastula stage embryos 24 to 26 h after fertilization.

genes encoding transcription factors that are causally responsible for specification of the skeletogenic regulatory state. From the GRN analysis came the prediction of the double-negative gate shown in Fig. 1D (9), and we have now identified the predicted missing component of this gate: The *r of mic* gene of the GRN model is *hesC*. In retrospect, the exact match between the predicted behavior of *r of mic* and the observed behavior of *hesC* is remarkable. Both the unique pattern of expression of *hesC*, which is not reproduced by any other known gene in this embryo, and the unique effects of preventing its expression are those required by the double-negative gate model in Fig. 1D. No additional players are likely to be inserted in the specification gate of Fig. 1D because manipulation of either component, *pmar1* overexpression or *hesC* underexpression, suffices to transform the whole embryo into cells specified as skeletogenic mesenchyme. In either perturbation, all cells (i) express the regulatory state of the skeletogenic micromere lineage (i.e., they transcribe the *delta*, *alx1*, *tbr*, and *ets1* genes, normally at this stage specific to the skeletogenic micromere lineage; in the *pmar1* overexpression, they even express terminal differentiation genes, such as *sm50*, not examined here); (ii) ingress into the blastocoel; and (iii) assume a mesenchymal form (Figs. 4 and 5 and SI Table 2) (9, 10).

Although this remains to be finally authenticated by identification of the *cis*-regulatory target sites, HesC interactions with the target genes of Fig. 1D are likely to be direct, as is likely to be the interaction of Pmar1 with the *hesC* regulatory apparatus. First, both genes encode proteins that contain transcriptional repression domains (see Results for HesC and ref. 9 for Pmar1). Second, the

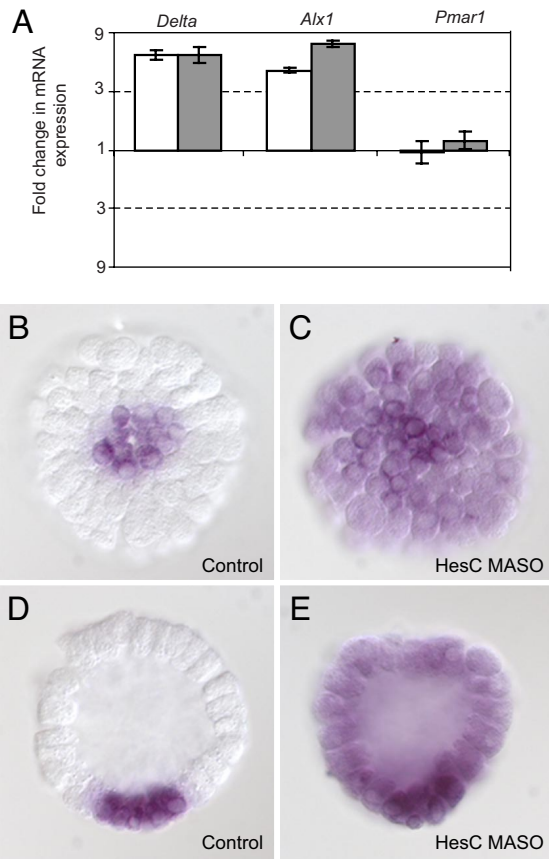


Fig. 5. Effect of HesC MASO on micromere lineage genes. (A) Fold change in *delta*, *alx1*, and *pmar1* mRNA expression in HesC MASO embryos relative to unperturbed embryos (12 h after fertilization). Fold change representation is as in Fig. 2. Open and filled bars indicate two independent batches of embryos. (B–E) Images of embryos (12 h after fertilization) on which WMISH was performed by using *delta* probe. Vegetal view (B) and side view (D) of an unperturbed embryo and vegetal view (C) and side view (E) of an HesC MASO embryo are shown.

kinetics with which the gate operates almost precludes any intervening steps. In sea urchin embryos at 15°C, it requires ≈ 2 –3 h for a regulatory gene to be activated, its product to be translated and transported to the nucleus, and a target gene to respond (20). We show here (Fig. 3A) that zygotic expression of *hesC* starts only ≈ 2 h after that of *pmar1*, and the zygotic expression of *delta* starts only ≈ 2 h after that of *hesC*. Third, we have carried out a *cis*-regulatory analysis of the *delta* gene, the key relevant results of which are summarized in SI Fig. 7. For the present discussion, the most important findings are: (i) the *cis*-regulatory module controlling *delta* expression in the micromere lineage responds to *pmar1* overexpression, just as does the endogenous *delta* gene (i.e., by ubiquitous ectopic expression), and also responds in just the same way to *hesC* MASO, again like the endogenous gene; however (ii), a region can be isolated from it that causes ubiquitous expression in unperturbed embryos and is not subject to repression, whereas other regions repress the ectopic expression and are subject to *pmar1* control. These results exclude the possibility that there is an indirect effect such that HesC represses another gene that is in turn responsible for *delta* activation because then the *cis*-regulatory DNA regions responsible for activation and repression would not be physically separable. Because HesC has a WRPW sequence, like all of its orthologues, it must act as a repressor, and we have shown that it does not interfere with *pmar1* expression. These observations strongly indicate repressive interaction at the *delta* *cis*-regulatory module.

The Double-Negative Gate. The main feature of this mechanism is the use of two repressors in regulatory tandem, and nonlocalized activators to produce a highly confined spatial pattern of gene expression. This mechanism is not uncommon; for example, the dorsal–ventral GRN for the early *Drosophila* embryo (21) affords several examples that are in essence similar. An alternative first step would be the highly localized expression of activators. This mechanism is common in later development, but in the early embryo, the boundaries of expression domains are often controlled negatively by the activation of repressors (1). In the sea urchin embryo, the known maternal regulatory transcripts are all globally distributed (12, 15, 16). Early on, before territorial regulatory states have been established, regional activation of repressors in response to initial anisometric cues is as parsimonious a strategy as regional activation of activators (1). An additional advantage of the double-negative gate is that it provides, de facto, the active repression of regulatory states outside the correct domain of their expression. Thus, it acts as an exclusion effect (22), actively ensuring silence of target genes in ectopic locations while ensuring their expression in correct locations.

Evolutionary Implications. Sea urchins are the only echinoderm class that produces an embryo/larva skeleton from a precociously specified micromere lineage. Thus, the regulatory apparatus for skeletogenic micromere specification, including the double-negative gate, arose in this lineage. An idea proposed earlier is that generation of the larval skeleton evolved as a cooption of the gene regulatory program for the production of the adult calcite skeleton (9, 23). The *hesC*–*pmar1* double-negative gate provides in principle a particularly economical means for highjacking the downstream skeletogenic regulatory machinery. Part of the circuitry is likely to have been already available. The Hairy/E(spl) family factors are used to repress the *delta* gene across the Bilateria (e.g., in both insect and vertebrate nervous system development) (24, 25). Sea urchin HesC repression of *delta* may indicate the inclusion in the cooption process of an ancient widespread plug-in (i.e., a conserved GRN linkage that is used in multiple, entirely unrelated, developmental contexts) (26). Now that the regulatory players are all in hand and most of their roles known, it should be possible to experimentally explore the evolution of the sea urchin skeletogenic specification by synthetically recreating the regulatory steps that led to its existence.

Materials and Methods

Animals, *pmar1* MOE, and *hesC* MASO. *pmar1* was overexpressed (i.e., its expression was forced in all cells of the embryos) by microinjecting *pmar1* mRNA into fertilized eggs. Microinjection solutions were prepared containing 25 ng/μl of *pmar1* mRNA and 0.12 M KCl.

Translation of HesC transcripts was blocked by microinjection of *hesC* MASO into fertilized eggs. MASO was synthesized (Gene Tools, Philomath, OR) complementary to the sequence of the first 25 bp of the coding region of *hesC*. The sequence of the oligonucleotide was: 5′-GTTGGTATCCAGATGAAGTAAGCAT-3′. Microinjection solutions were prepared containing 0.12 M KCl and 100, 250, or 500 μM *hesC* MASO.

Gametes from *S. purpuratus* were microinjected as described (27). We aimed at microinjecting a volume of ≈10 pl. Unperturbed embryos from the same batch were used as a control. Living embryos were visualized at chosen developmental time points on an AxioScope 2 Plus microscope (Carl Zeiss, Hall-

bergmoos, Germany) equipped with the recording device Axio-Cam MRm (Carl Zeiss).

Quantification of mRNA. An RNeasy Micro Kit (74004; Qiagen, Valencia, CA) was used to isolate RNA from samples of ≈100 embryos as described in the manufacturer's manual. cDNA was prepared from these samples by RT-PCR. The iScript cDNA Synthesis Kit (170–8891; Bio-Rad, Hercules, CA) was used for this purpose.

Quantitative PCR was conducted as described (27) by using primer sets designed to produce amplicons of 125 to 150 bp (for primer sequences, see <http://supg.caltech.edu/resources/methods/q-pcr.psp>). Amplification reactions were analyzed on an ABI 7900HT Fast Real-Time PCR System by using SYBR Green chemistry (iScript One-Step RT-PCR Kit; Bio-Rad). Levels of *ubiquitin* mRNA are known to remain relatively constant (≈220,000 molecules per embryo) during the relevant developmental stages (28, 29) and were used as an internal standard to determine the levels of mRNA per embryo of all other genes.

WMISH. Digoxigenin (DIG)-labeled RNA probes were prepared as described (30). A DIG-labeled HesC probe was transcribed from the HesC cDNA clone yde51c06 (CX199264; from a *S. purpuratus* EST library) kindly provided by James Coffman (Mount Desert Island Biological Laboratory, Salisbury Cove, ME). A sense DIG-labeled control probe was transcribed from the same clone, which does not recognize any known or predicted transcript. Dinitrophenol (DNP)-labeled RNA *delta* probe was prepared as described (31) by using the same plasmid as used for the DIG-labeled *delta* probe of ref. 9.

WMISH was performed by using a standard method, as described in refs. 32 and 33, with minor modifications (Sagar Damle and E.H.D., unpublished data). Hybridization reaction and washes were carried out at 65°C. Concentration of probe in hybridization reaction was 1 ng/μl. Antibody incubation was carried out containing a 1,000-fold dilution of anti-DIG antibody (Fab fragments; Roche Diagnostics, Indianapolis, IN) for DIG-labeled probes or anti-DNP antibody-alkaline phosphatase (Mirus, Madison, WI) for DNP-labeled probes.

Double-WMISH protocol (from Sagar Damle and E.H.D., unpublished data) was based on the above protocol for WMISH and the double-WMISH protocol described earlier (34). Steps before the hybridization reaction were as described above for WMISH. The hybridization reaction was carried out containing two probes (1 ng/μl each): a DNP-labeled probe and a DIG-labeled probe. Anti-DIG antibody was used for the first antibody incubation. The first staining reaction (purple) was then carried out as described above for the WMISH protocol, with NBT (N-6876; Sigma-Aldrich, St. Louis, MO)/BCIP (B-8503; Sigma-Aldrich). The staining reaction and antibody activity were stopped as in ref. 34. Anti-DNP antibody-alkaline phosphatase was used in the second antibody incubation. The second staining reaction (orange) was similar to the first one, except that INT (1–8377; Sigma-Aldrich)/BCIP was used instead of NBT/BCIP.

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- Davidson EH (2006) *The Regulatory Genome: Gene Regulatory Networks In Development and Evolution* (Academic, New York).
- Davidson EH, Rast JP, Oliveri P, Ransick A, Calestani C, Yuh CH, Minokawa T, Amore G, Hinman V, Arenas-Mena C, et al. (2002) *Science* 295:1669–1678.
- Oliveri P, Davidson EH (2004) *Curr Opin Genet Dev* 14:351–360.
- Levine M, Davidson EH (2005) *Proc Natl Acad Sci USA* 102:4936–4942.

- McClay D, Peterson R, Range R, Winter-Vann A, Ferkowicz M (2000) *Development (Cambridge, UK)* 127:5113–5122.
- Sweet H, Hodor P, Etensohn C (1999) *Development (Cambridge, UK)* 126:5255–5265.
- Sweet HC, Gehring M, Etensohn CA (2002) *Development (Cambridge, UK)* 129:1945–1955.

8. Livingston BT, Killian C, Wilt F, Cameron A, Landrum M, Ermolaeva O, Sapojnikov V, Maglott D, Buchanan A, Etensohn C (2006) *Dev Biol* 300:335–348.
9. Oliveri P, Carrick DM, Davidson EH (2002) *Dev Biol* 246:209–228.
10. Etensohn CA, Illies MR, Oliveri P, De Jong DL (2003) *Development (Cambridge, UK)* 130:2917–2928.
11. Howard-Ashby M, Materna SC, Brown CT, Tu Q, Oliveri P, Cameron RA, Davidson EH (2006) *Dev Biol* 300:27–34.
12. Materna SC, Howard-Ashby M, Gray RF, Davidson EH (2006) *Dev Biol* 300:108–120.
13. Rizzo F, Fernandez-Serra M, Squarzone P, Archimandritis A, Arnone MI (2006) *Dev Biol* 300:35–48.
14. Tu Q, Brown CT, Davidson EH, Oliveri P (2006) *Dev Biol* 300:49–62.
15. Howard-Ashby M, Materna SC, Brown CT, Chen L, Cameron RA, Davidson EH (2006) *Dev Biol* 300:74–89.
16. Howard-Ashby M, Materna SC, Brown CT, Chen L, Cameron RA, Davidson EH (2006) *Dev Biol* 300:90–107.
17. Kageyama R, Ohtsuka T, Hatakeyama J, Ohsawa R (2005) *Exp Cell Res* 306:343–348.
18. Paroush Z, Finley RL, Kidd T, Wainwright SM, Ingham PW, Brent R, Ish-Horowitz D (1994) *Cell* 79:805–815.
19. Grbavec D, Stifani S (1996) *Biochem Biophys Res Commun* 223:701–705.
20. Bolouri H, Davidson EH (2003) *Proc Natl Acad Sci USA* 100:9371–9376.
21. Stathopoulos A, Levine M (2005) *Dev Biol* 280:482–493.
22. Oliveri P, Davidson EH (2007) *Science* 315:1510–1511.
23. Hinman V, Nguyen A, Cameron RA, Davidson EH (2003) *Proc Natl Acad Sci USA* 100:13356–13361.
24. Beatus P, Lendahl U (1998) *J Neurosci Res* 54:125–136.
25. Gibert J, Simpson P (2003) *Int J Dev Biol* 47:643–651.
26. Davidson EH, Erwin DH (2006) *Science* 311:796–800.
27. Rast JP, Amore G, Calestani C, Livi CB, Ransick A, Davidson EH (2000) *Dev Biol* 228:270–286.
28. Nemer M, Rondinelli E, Infante D, Infante A (1991) *Dev Biol* 145:255–265.
29. Ransick A, Rast JP, Minokawa T, Calestani C, Davidson EH (2002) *Dev Biol* 246:132–147.
30. Minokawa T, Rast JP, Arenas-Mena C, Franco CB, Davidson EH (2004) *Gene Expression Patterns* 4:449–456.
31. Long S, Rebagliati M (2002) *BioTechniques* 32:494–498.
32. Croce J, Lhomond G, Gache C (2003) *Mech Dev* 120:561–572.
33. Bradham CA, McClay DR (2006) *Development (Cambridge, UK)* 133:21–32.
34. Minokawa T, Wikramanayake A, Davidson EH (2005) *Dev Biol* 288:545–558.