Regulatory Network of the Scoliosis-Associated Genes Establishes Rostrocaudal Patterning of Somites in Zebrafish

HIGHLIGHTS
- A core network establishes rostrocaudal polarity of segmented somites in zebrafish
- mesp genes link the segmentation clock with the FGF signaling gradient
- Gradual patterning is done by the action of different genes at different positions

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Article

Regulatory Network of the Scoliosis-Associated Genes Establishes Rostrocaudal Patterning of Somites in Zebrafish

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SUMMARY

Gene regulatory networks govern pattern formation and differentiation during embryonic development. Segmentation of somites, precursors of the vertebral column among other tissues, is jointly controlled by temporal signals from the segmentation clock and spatial signals from morphogen gradients. To explore how these temporal and spatial signals are integrated, we combined time-controlled genetic perturbation experiments with computational modeling to reconstruct the core segmentation network in zebrafish. We found that Mesp family transcription factors link the temporal information of the segmentation clock with the spatial action of the fibroblast growth factor signaling gradient to establish rostrocaudal (head to tail) polarity of segmented somites. We further showed that cells gradually commit to patterning by the action of different genes at different spatiotemporal positions. Our study provides a blueprint of the zebrafish segmentation network, which includes evolutionarily conserved genes that are associated with the birth defect congenital scoliosis in humans.

INTRODUCTION

A challenge in developmental biology is providing a molecular description of the cascade of regulatory steps that result in morphological changes and cell differentiation. Vertebrate somite segmentation is an example of a regulatory cascade and provides a system for studying the coordinated expression of multiple genes controlled by interconnected signaling pathways. Errors in this regulatory cascade result in various birth defects, including congenital scoliosis (Pourquie, 2011). Somites, the embryonic origin of the body segments of a vertebrate, are produced sequentially from the presomitic mesoderm (PSM) at the tail end of the embryo as the PSM elongates posteriorly (Figure 1A). In the zebrafish, one bilateral pair of somites segments at the anterior end of the PSM every 30 min. The timing of the separation of the somites from the PSM, referred to as somite periodicity, is controlled by a segmentation clock in the posterior PSM. This segmentation clock exhibits oscillatory expression of “clock” genes (Pourquie, 2011). The pacemaker mechanism of the segmentation clock relies on the auto-inhibitory feedback loop of her (in zebrafish) or Hes (in mouse) gene expression (Ay et al., 2013; Giudicelli et al., 2007; Harima et al., 2013; Jensen et al., 2003; Lewis, 2003; Monk, 2003; Schröter et al., 2012).

The positions of segment boundaries are instructed by the posteroanterior gradient of fibroblast growth factor (FGF) signaling (Simsek and Özbudak, 2018). The high FGF signal in the posterior region keeps the cell in an undifferentiated and developmentally plastic state in which the segmentation clock “ticks” (expression of clock genes oscillate) at a steady rapid rate (Pourquie, 2011). As the cell shifts into the anterior region of the PSM, the period of gene oscillations increases; consequently, the segmentation clock genes display dynamic expression patterns (stripes) in the PSM (Figure 1B). As the cells shift anteriorly and the oscillation period increases, the stripes of clock gene expression become narrower and become established in the posterior compartments of the prospective somite segments. In parallel, additional genes begin to be expressed dynamically in the anterior PSM. The expression of the genes becomes stable (zero or non-zero amounts) as the cell emerges from the PSM and becomes incorporated into a somite (Figure 1A). Different sets of genes are expressed in cells located in complementary (anterior or posterior) compartments of somites (rostrocaudal polarity), and thereby they govern the consequent differentiation of segmented cells (Holley, 2007; Stellabotte and Devoto, 2007). The final state of the cell depends on its phase of oscillation as it exits from the anterior end of PSM. In this way, the temporal oscillation in...
the posterior PSM is mapped into a periodic spatial pattern of cells in different states in the formed somites. Although the mechanism that generates the striped pattern of clock gene expression is well studied (Ay et al., 2013, 2014; Harima et al., 2013; Schroter et al., 2012), the regulatory network that establishes rostrocaudal somite polarity is not well characterized. The signaling pathways implicated in the establishment of rostrocaudal somite polarity include the FGF, Wnt, and Notch pathways (Pourquie, 2011).

Mutations of genes in the HES and MESP families of transcription factors result in scoliosis (Turnpenny et al., 2007), indicating their importance in somite development. Mesp genes encode transcription factors of the bHLH family, and Mesp2 deletion in mouse disrupts segmentation and rostrocaudal polarity of somites (Saga et al., 1997). In zebrafish, four Mesp2 homologs (mespaa, mespab, mespba, and mespbb) are
dynamically expressed in the anterior PSM, with peaks that coincide with troughs in the expression patterns of the clock genes her1 and her7 and the gene deltaC, encoding a ligand of the Notch pathway (Cutty et al., 2012; Sawada et al., 2000) (Figure 1B). In the PSM, mesp genes are expressed in the rostral compartment, whereas her and deltaC genes are expressed in the caudal compartment of the next segmenting somite. This complementarity is a prerequisite for the rostrocaudal polarity of segmented somites. Although rostrocaudal polarity is disrupted in zebrafish mesp mutants, mutation of the mesp family genes results in milder segmentation defects in zebrafish than in mouse owing to genetic and functional redundancy (Yabe et al., 2016). In zebrafish, the transcriptional repressor encoded by ripply1 is expressed in stripes in the anterior PSM and in recently formed somites (Kawamura et al., 2005). Morpholino-oligonucleotide-mediated knockdown of ripply1 in zebrafish or knockout of Ripply2 in mice causes segmentation defects (Kawamura et al., 2005; Morimoto et al., 2007). Mutations in MESP2 and RIPPLY2 are present in patients with congenital scoliosis (Cornier et al., 2008; Turnpenny et al., 2007) or Klippel-Feil syndrome (Karaca et al., 2015), respectively. Despite their developmental and medical importance, the regulatory relationships between the segmentation clock and the mesp genes are not completely understood. Joint knockdown, using morpholino-oligonucleotides, of her1 and her7 results in loss of rostrocaudally polarized expression of mesp genes in zebrafish (Henry et al., 2002; Oates and Ho, 2002). Whether this effect on the mesp gene expression pattern is due to a rapid action by Her family transcription factors or indirectly through Notch signaling is not clear.

To assess the regulatory relationships between the segmentation clock and morphogen signaling cascades in somite patterning, we have studied somite segmentation in zebrafish by altering the expression of specific genes in a time-controlled manner and by assessing the impact on the expression of other genes at defined times during development (Lewis and Ozbudak, 2007). Because cells at various stages of somitogenesis are distinctly positioned along the anteroposterior axis, global perturbation of gene expression abruptly at specific time during development will result in cells in different positions along the body axis experiencing the disturbance at different times relative to the time of their exit from the PSM. Thus the resulting somite segmentation pattern provides a map of the effects of the disturbance of gene expression relative to the time of exit from the PSM (Lewis and Ozbudak, 2007).

Here, we constructed a regulatory network that incorporates the zebrafish homologs of scoliosis-linked genes and provides a mechanism for the establishment of rostrocaudal segment polarity. We combined time-controlled perturbation experiments at high temporal resolution in zebrafish embryos with mathematical modeling to determine that rostrocaudally polarized expression of genes encoding the Mesp family of transcription factors is established by both the temporal action of the segmentation clock and the spatial action of FGF signaling gradient. We also showed that cells are gradually time stamped: their eventual state in the somite is dictated by the action of different transcription factors sequentially at different spatiotemporal positions during segmentation. Computational simulations of the segmentation network recapitulated the complementary expression of the families of her and mesp genes in wild-type embryos and the effects of perturbation experiments at the time window of 30 min to 4 h. This study provides a blueprint of the zebrafish segmentation network, incorporating genes with homologs in humans that are associated with congenital scoliosis.

RESULTS
Transcription of mespaa Is Rapidly Repressed by the Segmentation Clock

Owing to the complementary expression of her and mesp family genes, we hypothesized a regulatory interaction between them. To determine if mesp genes are regulated by segmentation clock transcription factors, we performed time-controlled perturbation experiments. We used the hsp70I:HA-her7 transgenic line to overexpress the her7 clock gene by heat shock. We compared the expression pattern of mespaa, mespba, and ripply1 in wild-type embryos and the hsp70I:HA-her7 transgenic embryos immediately after a 30-min heat shock and 30 or 60 min after the 30-min heat shock (Figure 2A). In situ hybridization (ISH) experiments in the embryos overexpressing her7 revealed reduced amounts of mespaa transcripts with their mRNAs barely detectable or undetectable immediately after the 30-min heat shock (Figure 2B). In contrast, mespba and ripply1 showed reduced and altered patterns of expression by 30 min after the heat shock period. Cells expressing high and low levels of mespba or ripply1 were intermingled by 60 min after the heat shock period (Figures 2C and 2D), reflecting desynchronization of expression patterns among neighboring cells (Ozbudak and Lewis, 2008). The reduction in mespaa in the hsp70I:HA-her7 transgenic lines within the 30-min heat shock treatment suggested a potential direct regulation of mespaa by
Figure 2. Expression of mesp Genes Read Out the Segmentation Clock Genes and FGF Signaling

(A) Embryos from different genetic backgrounds were fixed immediately or after 30 min, or 60 min of recovery following 30-min heat shock at 37°C.

(B–D) ISH images of mespaa (B), mespba (C), and rippy1 (D) after heat shock of tg(hsp70l:HA-her7) and wild-type (WT) embryos at different recovery time points. This experiment was repeated twice and 43–49 embryos analyzed for all probes and time points.

(E) Embryos from different genetic backgrounds were fixed after 30, 60, 90, 120, or 150 min of heat shock at 37°C.

(F) Flat mounted ISH images of mespba after heat shock of tg(hsp70l:dngfr1a-EGFP), tg(hsp70l:tc711a-GFP), and wild-type (WT) embryos at different time points. For each time point 8 to 22 embryos were analyzed. Red arrows show the interstripe distance, which was measured between the anterior ends of stripes.
We monitored somite boundaries by ISH for boundary establishment between the 9th and 10th somites in an embryo subjected to heat-shock-mediated mespab transgenic embryos were subjected to heat shock for 40 min; genes for segment boundary formation in zebrafish using transgenic lines (Windner et al., 2015) too v e r genes) (Yabe et al., 2016). We investigated the role of rostrocaudally polarized expression of mesp genes in zebrafish) and functional redundancy (action of yet-to-be discovered genes functionally equivalent to mesp genes) (Aulehla et al., 2008; Bajard et al., 2014; Delfini et al., 2005; Sawada et al., 2001). To elucidate which signal defines the spatial domain of mesp expression, we performed time-controlled perturbation experiments. As we have done previously (Simsek and Ozbudak, 2018), we used heat-shock-inducible transgenic lines to inhibit the activities of each signaling pathway in a time-controlled manner. We used hsp70l:cnfgr1a-EGFP, expressing dominant-negative FGF receptor fused with green fluorescent protein (GFP) reporter, to block FGF signaling (Lee et al., 2005) and hsp70l:tcfl1a-GFP, expressing dominant-negative tcfl1a fused with GFP reporter, to block Wnt-regulated transcriptional responses (Lewis et al., 2004). We subjected transgenic or wild-type embryos to heat shock for various durations (30–150 min) and fixed the embryos right after heat shock (Figure 2E). We performed ISH against mespaa (Figure S1) and mespba transcripts (Figure 2F) and measured the distance between “stripes” of mespba expression and the width of the posterior mespba stripe (Figures 2G and 2H). We observed an immediate increase in the width of the posteriormost mespaa or mespba stripes after the 30-min inhibition of FGF signaling. We also observed an increase in the interstripe distance, resulting from a posterior shift in the expression stripe, in the next segmentation cycle (60 min heat shock). In contrast, inhibition of Wnt signaling increased the width of the mespaa and mespba stripes only after a long delay after heat shock (Figures 2F–2H and S1). These data suggested a more direct effect of FGF signaling on mesp expression and an indirect effect of Wnt signaling. The striped pattern of expression is consistent with mesp genes responding to the temporal information from the clock with FGF providing spatial regulation of expression.

**Different Transcription Factors Function at Different Positions to Control Segmentation**

Mesp genes are expressed in only the anterior portions of prospective somites (Cutty et al., 2012; Morimoto et al., 2005; Sawada et al., 2000). Studies in mice indicated that the juxtaposition of cells expressing Mesp genes with those not expressing them (rostrocaudally polarized expression) is a requirement for the formation of the segment boundary in the last-forming somite (Morimoto et al., 2005). Mutations of mesp family genes result in segmentation defects in zebrafish that are less severe than those that result in mice. This phenotypic difference is attributed to both genetic redundancy (presence of duplicated mesp genes in zebrafish) and functional redundancy (action of yet-to-be discovered genes functionally equivalent to mesp genes) (Yabe et al., 2016). We investigated the role of rostrocaudally polarized expression of mesp genes for segment boundary formation in zebrafish using transgenic lines (Windner et al., 2015) to overexpress mespab and mespbb in a time-controlled manner during somitogenesis.

Heat-shock-driven expression of mespab or mespbb for 30 min or her7 for 40 min did not disrupt the next 1 to 3 somites that formed but resulted in defects in somites that formed later (Figures 2I, and 2J). We monitored somite boundaries by ISH for xirp2 (Figure 2I). For example, we observed a defect in somite boundary establishment between the 9th and 10th somites in an embryo subjected to heat-shock-mediated induction of her7 at the 4-somite stage (Figure 2I). The delay in the onset of the boundary defect decreased as the genes were induced later during somitogenesis (Figure 2J). These results showed that rostrocaudally...
polarized expression of mesp genes is not required for the next forming somite but is required for somites that form more posteriorly. Because segmentation defects started later in her7 overexpression than mespab or mespbb overexpression at all stages (Figure 2J), these results suggested that rostrocaudally polarized expression of mesp genes is necessary at a later time point (at more anterior positions) than the action of the segmentation clock (Figures 2I and 2J). Collectively, these results suggested that mesp genes function in the transfer of temporal information from the segmentation clock to the spatial patterning of somite segments. Furthermore, genes in the her and mesp families function at different times and positions after which cells no longer need their action to execute segmental pattern formation.

**Notch Signaling Activates Rostrocaudally Expressed mesp and her Genes with Different Kinetics**

Based on mutant and transgenic phenotypes in mouse, it was proposed that Notch signaling activated by the ligand Delta is an output of the segmentation clock, and Notch signaling drives striped expression of Mesp2 in the anterior PSM and thereby establishes anteroposterior compartmentalization of somites (Oginuma et al., 2010). According to this model, overactivation or inhibition of Notch signaling should disturb rostrocaudally polarized expression of Mesp genes and rapidly produce segmentation defects. This model is inconsistent with the following published data: (1) zebrafish and mouse embryos with hyperactive Notch signaling throughout the PSM still form more than 10 somites (Feller et al., 2008; Ozbudak and Lewis, 2008) and (2) segmentation proceeds for many cycles when Notch signaling is inhibited in both zebrafish and mouse (Huppert et al., 2005; Mara et al., 2007; Ozbudak and Lewis, 2008; Pourquier, 2011; Riedel-Kruse et al., 2007).

To clarify the role of Delta/Notch signaling, we blocked this pathway by treating embryos with γ-secretase inhibitor, N-[N-(3,5-difluorophenacetyl)-L-alanyl]-S-phenylglycine t-butyl ester (DAPT) during somitogenesis in a time-controlled manner (Figure 3A). The results showed that expression of mespaa, but not mespba or or her7, became barely detectable within 2 h of inhibition of Notch signaling (Figures 3B–3D). Consistent with previous work (Ozbudak and Lewis, 2008), the pattern of her7 expression was disrupted after 4 h (Figure 3D). Disruption in mespaa expression occurred after 4 h (Figure 3C). These results suggested that expression of mespaa had a stronger dependence on Notch signaling than does the expression of mespba and her7, because mespba and her7 expression became gradually desynchronized over a prolonged time period when Notch signaling is blocked and mespaa expression was affected more quickly. Redundancy in the mesp genes and the relatively low effect that we observed on mespba compared with mespaa in response to loss of Notch signaling may explain why segmentation proceeds normally over a long time when Notch signaling is disrupted. Therefore our data support a model in which Notch signaling primarily functions upstream of the segmentation clock in the posterior end of PSM and not downstream of the segmentation clock in the anterior PSM, as was proposed previously (Oginuma et al., 2010).

**Scoliosis-Linked Genes Are Connected in a Regulatory Network**

Mutations in Delta, Mesp, and Hes genes result in scoliosis in patients (Giampietro et al., 2009; Pourquie, 2011), and mutations of their orthologous genes in the mouse (zebrafish) model completely (partially) recapitulate the phenotype (Lleras Forero et al., 2018; Wopat et al., 2018). Although the functional importance of these genes has been better established in the mouse model (Morimoto et al., 2005; Oginuma et al., 2010), the dynamic regulatory relationships among these genes have not been determined in any model organism. To identify the dynamic regulatory relationships among delta, mesp, and hes genes, we overexpressed mespab and mespbb genes in a time-controlled manner and assessed their impact on the expression of her7, deltaC, mespaa, and mespba by performing ISH at different time intervals. For these experiments, we used a 60-min heat shock (Figure 3E). Expression of her7 was not affected by overexpression of either mesp genes, suggesting that there is no feedback loop between the clock and mesp genes (Figures 3F and 3G). Expression of deltaC was also not affected by overexpression of either mesp genes, suggesting that there is no feedback loop between the Delta/Notch signaling and mesp genes (Figures 3F and 3G). The genes mespaa and mespab are paralogs, as are mespba and mespbb. We used transgenic lines to overexpress one paralog and ISH probes to detect the changes in the expression levels of the other paralog gene. Overexpression of either of the mespab or mespbb genes reduced the expression of mespba, but not of mespaa (Figures 3H and 3I). However, transcription of mespba is reduced with 60-min heat shock (Figures 3H and 3I), but not 30-min heat shock (data not shown), suggesting that these regulations are indirect. These results are consistent with (and extend) the previous finding that transcription of mespb is reduced by an indirect negative feedback loop from Mespba that includes Ripply1 and Tbx6 (Takahashi et al., 2010; Windner et al., 2015).
To assess the regulatory relationship between mesp genes and ripply1, we used transgenic lines to overexpress mesp genes or ripply1 in a time-controlled manner (Figure 4A). Overexpression of either mespab or mespbb increased the transcription of ripply1 immediately after the 60-min heat shock (Figures 4B and 4C), whereas overexpression of ripply1 decreased transcription of both mespba and mespaa (Figures 4D and 4E). We combined our data with previous studies of the regulatory connections between mespba and ripply1 (Takahashi et al., 2010; Windner et al., 2015).
et al., 2015), and tbx6 and mespaa and mespba (Sawada et al., 2000; Windner et al., 2015) to construct a gene regulatory network (Figure 4F). The combined information suggested that mespb and ripply1 generate a negative feedback loop: mespb family proteins activate transcription of ripply1 and Ripply1 reduces the abundance of Tbx6, which is a transcriptional activator of mespb genes. We found that transcription of mespaa was not affected by the transient overexpression of mespab or mespbb (Figures 3H and 3I), suggesting that either Tbx6 was not eliminated by transient mesp overexpression or that transcription of mespba is more sensitive to Tbx6 levels than is mespaa transcription. In contrast, transcription of mespaa depends more strongly on Notch signaling than does transcription of mespba (Figures 3B and 3C). Thus we indicated regulation of the two mesp genes with different weights by Notch signaling and Tbx6 in our network of somitogenesis controlled by scoliosis-linked genes with the input of the segmentation clock and its intrinsic feedback as an autoinhibitory loop at the level of her (Figure 4F).

The regulatory relationships between the clock (her) and Notch (Figure 4F) are based on previous studies, indicating that the main function of Notch signaling is to synchronize oscillations of her and hes genes in the
posterior PSM (Delaneu et al., 2012; Mara et al., 2007; Ozbudak and Lewis, 2008; Riedel-Kruse et al., 2007), and our data, showing that her7 depends on Notch signaling (Figure 3D). However, this dependence is delayed relative to the dependence of mespa expression on Notch signaling (Figure 3B). Thus our regulatory network is consistent with our time-controlled perturbation data in previous Notch gain and loss-of-function experiments in zebrafish and mouse (Feller et al., 2008; Huppert et al., 2005; Ozbudak and Lewis, 2008; Pourquie, 2011). In the regulatory network (Figure 4F), Notch signaling regulates transcription of mespa by an incoherent feedforward loop: Notch activates transcription of mespa independently from the segmentation clock (her) and indirectly represses mespa transcription through Notch-mediated stimulation of the her family genes.

**Computational Modeling of the Segmentation Network Reproduces Complementary Expression of her and mesp Family Genes**

We built a data-driven mechanistic computational model of the segmentation network (Supplemental Information) to assess whether the regulatory network (Figure 4F) could recapitulate the complementary expression of mesp genes with that of her and delta in somitogenesis (Figure 1B). The model represented the tissue as a two-dimensional hexagonal lattice (4 by 50 cells) and simulated the abundance of her, delta, and mesp mRNAs, proteins, and protein complexes with time-delayed differential equations (see Transparent Methods for details). Biological phenomena are inherently noisy (stochastic) (Ozbudak et al., 2002). To reflect randomness in biochemical reactions of the network (Keskin et al., 2018), we performed pseudo-stochastic simulations of our model as previously described (Ay et al., 2014). Cell-to-cell variability in biochemical reactions was mimicked by assigning different rates for biochemical reactions in each cell (see Methods). This reaction rate variability was kept constant throughout the lifetimes of cells in the PSM (Ay et al., 2014). The simulations utilized previously measured (Ay et al., 2013, 2014; Giudicelli et al., 2007; Hanisch et al., 2013) or physiologically relevant reaction parameters (Table S1). To incorporate the repressive effect of FGF signaling on mesp transcription, the model restricts transcription of mesp genes only to anterior PSM cells.

We first constrained our model parameters by reproducing our earlier results (Ay et al., 2014). These simulations recapitulated previously published experimental data: (1) sustained, synchronized, and stripe-patterned oscillations of the segmentation clock genes in wild-type (Table S1) (Giudicelli et al., 2007); (2) notch1a−/− mutant phenotype where clock oscillation period increases (Herrgen et al., 2010), amplitude decreases (Ozbudak and Lewis, 2008), and oscillations desynchronize (Delaneu et al., 2012; Honkawa et al., 2006; Jiang et al., 2000; Mara et al., 2007; Ozbudak and Lewis, 2008; Riedel-Kruse et al., 2007) (Table S1); and (3) rapid repression of both her7 and deltaC genes after overexpression of her7 (Table S1 and Figure 5A) (Giudicelli et al., 2007).

We checked whether our model could reproduce the results of our time-controlled perturbation experiments. Using our model, we found parameter sets that reproduce our experimental data (Videos S1, S2, S3, and S4, and Table S2): (1) repression of mespa transcription by clock overexpression (model Figure 5A and Video S1, data Figure 2B), (2) gradual desynchronization of mespb transcription by clock overexpression (model Figure 5B and Video S2, data Figure 2C), (3) loss of mespa transcription when Notch signaling is blocked (model Figure 5C and Video S3, data Figure 3B), (4) gradual desynchronization of mespb transcription when Notch signaling is blocked (model Figures 5B and Video S4, data Figure 3C), and (5) differential dependence of mespb and mespa expression on overexpression of either gene with mespb repressed and mespa not affected (model Figure 5D, data Figures 3H and 3I). Note that although we did not impose any spatial restriction on the transcription of mesp genes in the anterior PSM, our model readily reproduced complementary transcription of her and mespb genes in wild-type embryos (Figure 5E and Videos S5 and S6). We further tested our model by showing that the striped expression of mesp genes are lost in her mutants (Video S7) demonstrating that her genes drive anteroposterior (A/P) polarized expression of mesp genes. However, repression of mespa by her genes alone cannot completely accommodate for the complementary expression of mesp and her genes because (1) deltaC is also repressed by her genes but is expressed concurrently with her genes (Giudicelli et al., 2007) and (2) mespb is not repressed by her genes (Figure 2C), but similar to mespa, it is also expressed in a complementary manner with her genes (Cutty et al., 2012; Sawada et al., 2000). Other feedback and feedforward loops in the network (Figure 4F) also contribute to the final dynamic expression patterns, and time-controlled perturbations of these feedback loops also disrupt A/P polarized complementary expression patterns (Figures 2, 3, and 4). Altogether, our results showed that the regulatory network established in this study is sufficient to explain the dynamic
expression patterns in wild-type zebrafish embryos and the results of previously published data as well as our time-controlled perturbation experiments.

**DISCUSSION**

Experimental analysis coupled with mathematical modeling is a powerful way to understand complex biological systems. Vertebrate segmentation is an excellent example of such a system, because it involves codependent expression of multiple genes, which is regulated by dynamic signaling pathways. The establishment of rostrocaudal polarity is critical for understanding the etiology and potentially
preventing various types of vertebral malformations. Somitogenesis shows the importance of timing in embryonic development. Hence, experiments that generate well-controlled temporal perturbations enable the investigation of questions related to timing using somitogenesis as the model system. By applying this strategy, we identified a high-resolution time course of transcriptional changes (with time window of 30 min to 4 h) and built a time-resolved regulatory network that establishes rostrocaudal polarity of somites in zebrafish. To test the sufficiency of the regulatory network in explaining experimental results, we built a comprehensive computational model. Computational simulations successfully recapitulated experimental results. More importantly, the model was essential to show that the regulatory network that we identified is sufficient to produce complementary expression of her and mesp genes.

Mesp genes are expressed only in the anterior PSM but excluded from the posterior PSM. Their restricted expression to the anterior PSM had been attributed to posterior-anterior gradients of FGF or Wnt signaling (Aulehla et al., 2008; Bajard et al., 2014; Delfini et al., 2005; Sawada et al., 2001); however, evidence was missing as to which of these signaling pathways controls the expression domain of mesp genes. By performing time-controlled perturbation experiments, we for the first time showed that altering FGF signaling immediately shifted expression domains of mesp genes more posteriorly, whereas Wnt signaling only shifted expression after a long delay (Figures 2F–2H). These results extend our previous observation that FGF signaling instructs both the position of the determination front and the anteriorly restricted expression of mesp genes, whereas Wnt signaling acts permissively upstream of the FGF signal (Simsek and Ozbudak, 2018). Expression of mesp genes in the anterior PSM is restricted to the anterior halves of prospective somites. This rostrocaudally polarized expression of Mesp genes has previously been attributed to activation by Notch signaling in the anterior PSM (Oginuma et al., 2010). In mouse, the segmentation clock had been proposed to affect Mesp gene expression indirectly through oscillations of Notch signaling. Mutation studies in zebrafish could not clarify whether the Her family clock proteins could regulate mesp gene expression more rapidly than Notch signaling and whether this regulation was mediated by activation or repression. Here, we found that the segmentation clock represses the expression of mespaa. Furthermore, the expression levels and patterns of both mespa and mespb have responded faster upon overexpression of Her7 (Figures 2B and 2C) than inhibition of Notch signaling (Figures 3B and 3C). These results suggest that the segmentation clock does not regulate the expression of mesp genes via Notch signaling as proposed previously based on steady-state mutant data in mice. Altogether our results indicated that mesp genes integrate the spatial information from the FGF gradient (such that mesp genes are expressed only in the anterior PSM, but not in the posterior PSM) and the temporal information from the clock (such that mesp genes are expressed in rostrocaudally polarized stripes rather than ubiquitously in the anterior PSM). Our results indicated that Notch signaling functions primarily upstream of the segmentation clock in the posterior PSM. Notch signaling regulated the transcription of mespaa through an incoherent feedforward loop, and transcription of mespb depended more on Tbx6 than on Notch signaling. Cells complete different steps of tissue patterning sequentially by the action of different transcription factors at different spatial locations along the axis.

We here carried out a multidisciplinary approach to map the regulatory network controlling rostrocaudal polarity of somites in the zebrafish model. This blueprint network developed in zebrafish is consistent with many of the phenotypes in mouse (Feller et al., 2008; Huppert et al., 2005; Oginuma et al., 2010; Pourquie, 2011), suggesting that at least the core elements are evolutionarily conserved between fish and mammals. Confirmation of our regulatory network in mammals awaits time-controlled perturbation studies like those we used here. Our model could easily be adapted to mammals at such time that time-controlled perturbation experiments could be performed. We anticipate our interdisciplinary quantitative approach to be adapted to reconstruct regulatory networks governing other dynamic questions in embryonic development.

Limitations of the Study

The reduction in mespaa levels in the hsp70:HA–her7 transgenic lines within 30-min heat shock treatment suggested a potential direct regulation of mespaa by Her7, whereas the delay in the change in mespb and ripply1 transcripts suggested an indirect regulation of mespb and ripply1 by Her7 (Figures 2B–2D). We have to note that the proof of a direct or indirect transcriptional regulation requires a separate in-depth study of transcriptional binding sites in future.
METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental Information includes Transparent Methods, one figure, two tables, and seven videos and can be found with this article online at https://doi.org/10.1016/j.isci.2019.01.021.

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AUTHOR CONTRIBUTIONS
E.M.Ö. and A.A. designed the project, built the mathematical model, and supervised the project. S.K. and M.F.S. performed the experiments and analysis of data. A.A., H.T.V., and C.Y. wrote the computational code and executed simulations. S.K., M.F.S., H.T.V., C.Y., S.H.D., A.A., and E.M.Ö. wrote the manuscript.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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REFERENCES

Aulehla, A., Wiegrebe, W., Baubet, V., Wahl, M.B., Deng, C., Taketo, M., Lewandoski, M., and Pourquie, O. (2008). Nat. Cell Biol. 10, 186–193.

Ay, A., Holland, J., Sperlea, A., Devakannalai, G.S., Knierer, S., Sangervasi, S., Stevenson, A., and Ozbudak, E.M. (2014). Spatial gradients of protein-level time delays set the pace of the traveling segmentation clock waves. Development 141, 4158–4167.

Ay, A., Knierer, S., Sperlea, A., Holland, J., and Ozbudak, E.M. (2013). Short-lived her proteins drive robust synchronized oscillations in the zebrafish segmentation clock. Development 140, 3244–3253.

Bajard, L., Morelli, L.G., Ares, S., Peceaux, J., Julicher, F., and Oates, A.C. (2014). Wnt-regulated dynamics of positional information in zebrafish somitogenesis. Development 141, 1381–1391.

Cornier, A.S., Staehling-Hampton, K., Delventhal, K.M., Saga, Y., Caubet, J.P., Sasaki, N., Ellard, S., Young, E., Ramirez, N., Carlo, S.E., et al. (2008). Mutations in the MESP2 gene cause spondylothyroidic dysostosis/Jarcho-Levin syndrome. Am. J. Hum. Genet. 82, 1334–1341.

Cutty, S.J., Fior, R., Henriques, P.M., Saude, L., and Wardle, F.C. (2012). Identification and expression analysis of two novel members of the Mesp family in zebrafish. Int. J. Dev. Biol. 56, 285–294.

Deleu, E.A., Francois, P., Shih, N.P., and Amacher, S.L. (2012). Single-cell-resolution imaging of the impact of notch signaling and mitosis on segmentation clock dynamics. Dev. Cell 23, 995–1005.

Delfini, M.C., Dubrulle, J., Malapert, P., Chal, J., and Pourquie, O. (2005). Control of the segmentation process by graded MAPK/ERK activation in the chick embryo. Proc. Natl. Acad. Sci. U.S.A 102, 11343–11348.

Dubrulle, J., McGrew, M.J., and Pourquie, O. (2001). FGF signaling controls somite boundary position and regulates segmentation clock control of spatiotemporal Fox gene activation. Cell 106, 219–232.

Feller, J., Schneider, A., Schuster-Gossler, K., and Gossler, A. (2008). Noncyclic Notch activity in the presomitic mesoderm demonstrates uncoupling of somite compartmentalization and boundary formation. Genes Dev. 22, 2166–2171.

Giampietro, P.F., Dunwoodie, S.L., Kusumi, K., Pourquie, O., Tassy, O., Offiah, A.C., Cornier, A.S., Alman, B.A., Blank, R.D., Raggio, C.L., et al. (2009). Progress in the understanding of the genetic etiology of vertebral segmentation disorders in humans. Ann. N. Y. Acad. Sci. 1151, 36–67.

Giudicelli, F., Ozbudak, E.M., Wright, G.J., and Lewis, J. (2007). Setting the tempo of the segmentation clock. Development 134, 1244–1253.

Holley, S.A. (2007). The genetics and embryology of zebrafish metamericism. Dev. Dyn. 236, 1422–1449.

Horikawa, K., Ishimatsu, K., Yoshimoto, E., Kondo, S., and Takeda, H. (2006). Noise-resistant and synchronized oscillation of the segmentation clock. Science 311, 677–688.

Jensen, M.H., Sneppen, K., and Tiana, G. (2003). Sustained oscillations and time delays in gene
Rare variants in the notch signaling pathway

H., Pehlivan, D., Jhangiani, S.N., Akdemir, Z.C., Karaca, E., Yuregir, O.O., Bozdogan, S.T., Aslan, 176–177.

signalling and the synchronization of the somite segmentation clock. Nature 408, 475–479.

Karaca, E., Yuregir, O.O., Bozdogan, S.T., Aslan, H., Pehlivan, D., Jhangiani, S.N., Akdemir, Z.C., Gambuän, T., Bayram, Y., Atik, M.M., et al. (2015). Rare variants in the notch signaling pathway describe a novel type of autosomal recessive Klippel-Feil syndrome. Am. J. Med. Genet. A 167A, 2795–2799.

Kawamura, A., Koshida, S., Hjikata, H., Ohbayashi, A., Kondoh, H., and Takada, S. (2005). Groucho-associated transcriptional repressor ripply1 is required for proper transition from the presomitic mesoderm to somites. Dev. Cell 7, 735–744.

Keskin, S., Devakannalai, G.S., Kwon, S.B., Vu, H.T., Hong, Q., Lee, Y.Y., Soltani, M., Singh, A., Ay, A., and Ozbudak, E.M. (2018). Noise in the vertebrate segmentation clock is boosted by time delays but tamed by notch signaling. Cell Rep. 23, 2175–2185.e4.

Lee, Y., Grill, S., Sanchez, A., Murphy-Ryan, M., and Poss, K.D. (2005). Fgf signaling instructs position-dependent growth rate during zebrafish fin regeneration. Development 132, 5173–5183.

Lewis, J. (2003). Autoinhibition with transcriptional delay: a simple mechanism for the zebrafish somitogenesis oscillator. Curr. Biol. 13, 1398–1408.

Lewis, J., and Ozbudak, E.M. (2007). Deciphering the somite segmentation clock: beyond mutants and morphants. Dev. Dyn. 236, 1410–1415.

Lewis, J.L., Bonner, J., Modrell, M., Ragland, J.W., Moon, R.T., Dorsky, R.I., and Raible, D.W. (2004). Reiterated Wnt signaling during zebrafish neural crest development. Development 131, 1299–1308.

Lleras Forero, L., Narayanan, R., Huitema, L.F., VanBergen, M., Apschner, A., Peterson-Maduro, J., Logster, I., Valentín, G., Morelli, L.G., Oates, A.C., et al. (2018). Segmentation of the zebrafish axial skeleton relies on notochord sheath cells and not on the segmentation clock. Elife 7, 1–28.

Mara, A., Schroeder, J., Chalouni, C., and Holley, S.A. (2007). Priming, initiation and synchronization of the segmentation clock by deltaD and deltaC. Nat. Cell Biol. 9, 523–530.

Monk, N.A. (2003). Oscillatory expression of Hes1, p53, and NF-kappaB driven by transcriptional time delays. Curr. Biol. 13, 1409–1413.

Mormito, M., Sasaki, N., Oginuma, M., Kiso, M., Igarashi, K., Aizaki, K., Kanno, J., and Saga, Y. (2007). The negative regulation of Mesp2 by mouse Ripply2 is required to establish the rostro-caudal patterning within a somite. Development 134, 1561–1569.

Mormito, M., Takahashi, Y., Endo, M., and Saga, Y. (2005). The Mesp2 transcription factor establishes segmental borders by suppressing Notch activity. Nature 435, 354–359.

Oates, A.C., and Ho, R.K. (2002). Hairy/E(spl)-related (Her) genes are central components of the segmentation oscillator and display redundancy with the Delta/Notch signaling pathway in the formation of anterior segmental boundaries in the zebrafish. Development 129, 2929–2946.

Oginuma, M., Takahashi, Y., Kizaiini, S., Kiso, M., Kanno, J., Kimura, A., and Saga, Y. (2010). The oscillation of Notch activation, but not its boundary, is required for somite border formation and rostral-caudal patterning within a somite. Development 137, 1515–1522.

Ozbudak, E.M., and Lewis, J. (2009). Notch signalling synchronizes the zebrafish segmentation clock but is not needed to create somite boundaries. PLoS Genet. 4, e15.

Ozbudak, E.M., Thattai, M., Kurtser, I., Grossman, A.D., and van Oudenaarden, A. (2002). Regulation of noise in the expression of a single gene. Nat. Genet. 31, 69–73.

Pourquie, O. (2011). Vertebrate segmentation: from cyclic gene networks to sclerotosis. Cell 145, 650–663.

Riedel-Kruse, I.H., Muller, C., and Oates, A.C. (2007). Synchrony dynamics during initiation, failure, and rescue of the segmentation clock. Science 317, 1911–1915.

Saga, Y., Hata, N., Koseki, H., and Taketo, M.M. (1997). Mesp2: a novel mouse gene expressed in the presegmented mesoderm and essential for segmentation initiation. Genes Dev. 11, 1827–1839.

Sawada, A., Fritz, A., Jiang, Y., Yamamoto, A., Yamashu, K., Kuroiwa, A., Saga, Y., and Takada, H. (2000). Zebrafish Mesp family genes, mesp-a and mesp-b are segmentally expressed in the presomitic mesoderm, and Mesp-b confers the anterior identity to the developing somites. Development 127, 1691–1702.

Sawada, A., Shinya, M., Jiang, Y.J., Kawakami, A., Kuroiwa, A., and Takada, H. (2001). Fgf/MAPK signalling is a crucial positional cue in somite boundary formation. Development 128, 4873–4880.

Schoiter, C., Ares, S., Morelli, L.G., Isakova, A., Hens, K., Soroldoni, D., Gajewski, M., Julicher, F., Maeki, S.J., Deplancke, B., et al. (2012). Topology and dynamics of the zebrafish segmentation clock core circuit. PLoS Biol. 10, e1001364.

Simsek, M.F., and Ozbudak, E.M. (2018). Spatial fold change of FGF signaling encodes positional information for segmental determination in zebrafish. Cell Rep. 24, 66–78.e8.

Stellabotte, F., and Devoto, S.H. (2007). The telost dermatomyotome. Dev. Dyn. 236, 2432–2443.

Takahashi, J., Ohbayashi, A., Oginuma, M., Saito, D., Mochizuki, A., Saga, Y., and Takada, S. (2010). Analysis of Ripply1/2-deficient mouse embryos reveals a mechanism underlying the rostro-caudal patterning within a somite. Dev. Biol. 342, 134–145.

Tumpenny, P.D., Alman, B., Cornier, A.S., Giampietro, P.F., Offiah, A., Tassy, O., Pourquie, O., Kubuski, K., and Dunwoodie, S. (2007). Abnormal vertebral segmentation and the notch signaling pathway in man. Dev. Dyn. 236, 1456–1474.

Windner, S.E., Doris, R.A., Ferguson, C.M., Nelson, A.C., Valentín, G., Tan, H., Oates, A.C., Wardle, F.C., and Devoto, S.H. (2015). Tbx6, Mesp-b and Ripply1 regulate the onset of skeletal myogenesis in zebrafish. Development 142, 1159–1168.

Wopat, S., Bagwell, J., Sumigray, K.D., Dickson, A.L., Huitema, L.F.A., Poss, K.D., Schulte-Merker, S., and Bagnat, M. (2018). Spine patterning is guided by segmentation of the notochord sheath. Cell Rep. 22, 2026–2038.

Yabe, T., Hoshijima, K., Yamamoto, T., and Takada, S. (2016). Quadruple zebrafish mutant reveals different roles of Mesp genes in somite segmentation between mouse and zebrafish. Development 143, 2842–2852.
Supplemental Information

Regulatory Network of the Scoliosis-Associated Genes Establishes Rostrocaudal Patterning of Somites in Zebrafish

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Figure S1. Expression of *mespaa* genes reads out the FGF signaling, Related to Figure 2. Flat mounted ISH images of *mespaa* transcripts at different time points of heat shock of *tg(hsp70l:dnfgfr1a-EGFP)*, *tg(hsp70l:tcf7l1a-GFP)* and wild-type (WT) embryos) at 37 °C. 8 to 22 embryos were flat-mounted for each time points.
## Table S1. Description of Simulation Parameters, Related to Figure 5

| Parameter | Description of parameter | Range     | Set  |
|-----------|--------------------------|-----------|------|
| \(msh\)  | her mRNA synthesis rate  | \([67.2,69.3]\) | 67.4 |
| \(msma\) | mespa mRNA synthesis rate| \([30.1,40.3]\) | 40.2 |
| \(msmb\) | mespb mRNA synthesis rate| \([46.6,50.9]\) | 50.1 |
| \(msd\)  | delta mRNA synthesis    | \([31.6,33.7]\) | 32.6 |
| \(mdh\)  | her mRNA degradation     | 0.5       | 0.5  |
| \(mdma\) | mespa mRNA degradation rate| 0.1    | 0.1  |
| \(mdmb\) | mespb mRNA degradation rate| \([0.1,0.11]\) | 0.101 |
| \(mdd\)  | delta mRNA degradation rate| 0.5    | 0.5  |
| \(psh\)  | her protein synthesis rate| \([10.4,10.7]\) | 10.6 |
| \(psma\) | mespa protein synthesis rate| \([34.4,53.4]\) | 43   |
| \(psmb\) | mespb protein synthesis rate| \([19.1,21.5]\) | 21.5 |
| \(psd\)  | delta protein synthesis rate| \([26.4,26.6]\) | 26.5 |
| \(pdh\)  | her protein degradation rate| \([0.183,0.185]\) | 0.183 |
| \(pdm\)  | mespa protein degradation rate| \([0.155,0.273]\) | 0.157 |
| \(pdmb\) | mespb protein degradation rate| \([0.1,0.227]\) | 0.109 |
| \(pdd\)  | delta protein degradation rate| 0.5    | 0.5  |
| \(dahh\) | her-her dimer association rate| 0.0003 | 0.0003 |
| \(damam\) | mespa-mespa dimer association rate| \([0.00293,0.00387]\) | 0.00297 |
| \(damamb\) | mespa-mespb dimer association rate| \([0.0256,0.03]\) | 0.029 |
| \(ddmbm\) | mespb-mespb dimer association rate| \([0.00337,0.00617]\) | 0.00381 |
| \(ddhh\) | her-her dimer dissociation rate| \([0.194,0.209]\) | 0.202 |
| \(ddmam\) | mespa-mespa dimer dissociation rate| \([0.00303,0.0252]\) | 0.00497 |
| \(ddmamb\) | mespa-mespb dimer dissociation rate| \([0.214,0.276]\) | 0.27 |
| \(ddmbmb\) | mespb-mespb dimer dissociation rate| \([0.0952,0.3]\) | 0.293 |
| \(pddh\) | her-her dimer degradation rate| \([0.17,0.172]\) | 0.171 |
| \(pdmam\) | mespa-mespa dimer degradation rate| \([0.179,0.222]\) | 0.202 |
| \(pdmamb\) | mespa-mespb dimer degradation rate| \([0.1,0.109]\) | 0.1 |
| \(pdmmbmb\) | mespb-mespb dimer degradation rate| \([0.438,0.489]\) | 0.439 |
| \(delaymh\) | her mRNA synthesis delay rate| \([7.7,0.01]\) | 7.01 |
| \(delaymm\) | mespa mRNA synthesis delay rate| 15     | 15   |
| \(delaymmmb\) | mespa mRNA synthesis delay rate| \([14.6,15]\) | 14.8 |
| \(delaymd\) | delta mRNA synthesis delay rate| \([8.99,9.05]\) | 9    |
| \(delayph\) | her protein synthesis delay rate| 1.14   | 1.14 |
| \(delaypa\) | mespa protein synthesis delay rate| \([0.4,0.524]\) | 0.407 |
| \(delaypb\) | mespb protein synthesis delay rate| \([0.4,0.47]\) | 0.405 |
| \(delaypd\) | delta protein synthesis delay rate| 12.6   | 12.6 |
| $\text{delta}_M$ | delay for indirect mesp-ripply-tbx6 feedback | [42.7,43.1] | 42.7 |
|-----------------|---------------------------------------------|-------------|------|
| $\text{critphh}$ | critical binding rate of her-her dimer       | [390,406]   | 394  |
| $\text{critipd}$ | critical binding rate of delta protein       | [603,651]   | 651  |
| $\text{critpm}_{a_m_a}$ | critical binding rate of mespa-mespa dimer | [1679,1986] | 1984 |
| $\text{critpm}_{b_m_b}$ | critical binding rate of mespb-mespb dimer  | [500,659]   | 501  |
| $\text{oehler}$  | her over expression rate                     | [27.8,31]   | 29.7 |
| $\text{oemespa}$ | mespa over expression rate                  | [30.9,96.3] | 53.5 |
| $\text{oemespb}$ | mespb over expression rate                  | [51.9,84.3] | 66.7 |
Table S2. Description of Simulation Conditions, Related to Figure 5

| Genetic Background | Tested Condition                                                                 | Check Time |
|--------------------|----------------------------------------------------------------------------------|------------|
| **Wildtype:**      | **Her** mRNA oscillation period is ~30 minutes.                                  | 100 – 300  |
|                    | **Her** mRNA expression shows sustained oscillation.                             | 100 – 300  |
|                    | **Her** mRNA period increases from posterior end of the PSM to the anterior end of the PSM. | 600 – 900  |
|                    | **Her** mRNA oscillations are synchronized between neighboring cells.             | 600 – 630  |
|                    | **Her** and mespa mRNAs show complementary pattern in the anterior PSM.          | 600 – 630  |
| **Notch1a^{−/−} mutant:** DeltaC protein synthesis is set to 0.                    | **Her** mRNA period increases 7%-20% in notch1a^{−/−} mutant.                    | 100 – 300  |
|                    | **Her** mRNA amplitude decreases 15%-70% in notch1a^{−/−} mutant.                | 600 – 630  |
|                    | **Her** mRNA oscillations are desynchronized in notch1a^{−/−} mutant.             | 600 – 630  |
|                    | **Mespa** mRNA amplitude decreases more than 70% in notch1a^{−/−} mutant.        | 600 – 630  |
|                    | **Mespb** mRNA oscillations are desynchronized in notch1a^{−/−} mutant.          | 600 – 630  |
| **Her overexpression:** Her protein synthesis is increased for 30 min starting after 600 min. | **Her** mRNA amplitude decreases more than 70% in 30 minutes after her overexpression. | 630 – 660  |
|                    | **DeltaC** mRNA amplitude decreases more than 70% in 30 minutes after her overexpression. | 630 – 660  |
|                    | **Mespa** mRNA amplitude decreases more than 70% in 30 minutes after her overexpression. | 630 – 660  |
|                    | **Mespb** mRNA oscillations are desynchronized in 1.5 hours after her overexpression. | 690 – 720  |
| **Notch signaling disruption by DAPT:** DeltaC protein synthesis is set to 0 after 600 min. | **Her** mRNA oscillations are desynchronized in 4 hours after DAPT treatment. | 840 – 870  |
|                    | **Her** mRNA amplitude decreases 15%-70% in 4 hours after DAPT treatment.         | 840 – 870  |
|                    | **Mespa** mRNA amplitude decreases more than 70% in 2 hours after DAPT treatment. | 720 – 750  |
|                    | **Mespb** mRNA oscillations are desynchronized in 4 hours after DAPT treatment.   | 840 – 870  |
| **Mespa overexpression:** Mespa protein synthesis is increased for 60 min starting after 600 min. | **Mespa** mRNA amplitude decreases more than 70% in 1 hour after mespa overexpression. | 660 – 690  |
| **Mespb overexpression:** Mespb protein synthesis is increased for 60 min starting after 600 min. | **Mespb** mRNA amplitude decreases more than 70% in 1 hour after mespb overexpression. | 660 – 690  |
TRANSPARENT METHODS

CONTACT FOR REAGENT AND RESOURCE SHARING
Further information and requests for reagents may be directed to the Lead Contact Ertugrul Ozbudak (Ertugrul.Ozbudak@cchmc.org).

EXPERIMENTAL MODEL AND SUBJECT DETAILS
Fish stocks
All the fish experiments were performed under the ethical guidelines of Albert Einstein College of Medicine and Cincinnati Children’s Hospital Medical Center, and animal protocols were reviewed and approved by the respective Institutional Animal Care and Use Committees (Protocol # 20150704 and Protocol # 2017-0048). Fish were kept on a 14-10 light/dark cycle at the Zebrafish Core Facility, maintained at 28.5°C. Transgenic lines hsp70:HA-her7 (Giudicelli et al., 2007), hsp70l:mespab-myc, hsp70l:mespbb-myc, hsp70l:ripply1-myc (Windner et al., 2015) hsp70l:dnfgr1a-EGFP (Lee et al., 2005) and hsp70l:tcf7l1a-GFP (Lewis et al., 2004) were used during this study.

METHOD DETAILS
Heat-shock procedures
We used heat-shock inducible promoters to perform time-controlled perturbation experiments throughout this study. This approach induces transgenes very rapidly (Giudicelli et al., 2007) as compared to alternative Tet-on inducible system (Watanabe et al., 2007; Wehner et al., 2015). Transgenic heterozygous fishes were crossed to wild-type fish to obtain transgenic and control embryos with equal proportions. Embryos were kept at a temperature range of 23-28°C until the desired stage for heat-shock. They were then transferred to pre-warmed E3 medium in a 37°C incubator for the desired length of time, then fixed immediately in ice-cold 4% paraformaldehyde or returned to 28°C for further development and fixation at a late recovery time point (Giudicelli et al., 2007). Temporal loss of function of Notch signaling was accomplished by exposing embryos to 100 µM of the γ-secretase inhibitor, N-[N-(3,5-difluorophenacetyl)-L-alanyl]-S-phenylglycine t-butyl ester (DAPT) in DMSO (Ozbudak and Lewis, 2008). Control embryos were exposed to DMSO.

In situ hybridization
In situ hybridization was performed according to standard protocols. Digoxigenin-labelled RNA probes were as previously described for her1 (Takke and Campos-Ortega, 1999), her7 (Henry et al., 2002; Oates and Ho, 2002), deltaC (Jiang et al., 2000) and cb1045 (xirp2a – Zebrafish Information Network) (Riedel-Kruse et al., 2007). Probes for mespaa, mespba, and ripply1 are generated using the nucleotides in between 1-887, 1-801, and 86-785, respectively. We validated the genotype of the selected embryos by PCR by using the following primers: mespabUprv: TCAACATTGGCATTTTTCAGG, mespabF_NOT1: gatGCgGcGCgC GCATTcaACTCAAGCTCCAGA, mespbbR_ECOR1: gatcGAATTCAGTGGACGCTTTTGTGTGA, mespbbF_NOT1: gatcGCGGCCTAGGGTGGTCTGGACAGG, shHsp70l_BbvClfw: gatcCCTCAGGCCACACAAACCGCACATTTTTC and ripply1rv: CCTCGACGTCACTTTTCA.

QUANTIFICATION AND STATISTICAL ANALYSIS
Mathematical model
We developed a delayed differential equation model (DDE) with 12 equations (see below) and 44 parameters (Table S1). Each equation in our DDE model represents the rate of change of a model state (mRNA, protein, or protein complex); each model parameter represents the rate of the corresponding reaction that influences the concentrations of the model states. Biological reaction terms describe the synthesis and degradation of mRNAs and proteins, as well as dimer association, dissociation, and degradation.

The genes included in the model are her, deltaC, mespa and mespb. Following Lewis (2003) (Lewis, 2003), we represented her1 and her7 genes as one her gene. In our model, Her protein forms Her-Her homodimer, and represses transcription of her, deltaC, and mespa genes (Figure 5E). Mespb proteins form dimers to repress transcription of mespb gene. DeltaC triggers the proteolytic cleavage of the Notch protein intracellular domain (NICD), NICD translocates into the nucleus and activates the transcription of her, mespa, and mespb. The transcriptional repressors Her-Her, Mespa-Mespa, and Mespb-Mespb compete with the NICD protein for binding to the DNA regulatory region to repress transcription of her, mespa and mespb genes (Ozbudak and Lewis, 2008). To simplify the model, we followed earlier work (Ay et al., 2014; Lewis, 2003) and did not explicitly write an equation representing the production of NICD. Instead, we represented NICD levels in each cell as a function of the DeltaC protein levels in all neighboring cells.

The variables: mh, md, mmh, and mmh represent the number of mRNA molecules of her, deltaC, mespa and mespb respectively; ph, pd, pmh and pmh represent the number of protein molecules of Her, DeltaC, Mespa and Mespb respectively; and phh, pmmh, pmmh and pmmh represent the number of molecules of Her-Her, Mespa-Mespa, Mespb-Mespb and Mespa-Mespb dimers, respectively. mRNA synthesis rates are denoted as msh, mdd, msh and msb for her, deltaC, mespa and mespb genes, respectively. mRNA degradation rates are denoted as mdh, mdd, mmh and mmdh for her, deltaC, mespa and mespb mRNAs, respectively. Protein synthesis rates are denoted as phh, psd, psmh and psmh for Her, DeltaC, Mespa and Mespb proteins, respectively. Degradation rates for Her, DeltaC, Mespa and Mespb proteins are denoted as pdh, pdd, pmdh and pmdh, respectively. Dimer association, dissociation, and degradation rates for Her-Her are represented by dahlh, ddhh and pdhh, respectively. Dimer association, dissociation, and degradation rates for Mespa-Mespa, Mespb-Mespb and Mespa-Mespb are represented by dmmmh, ddmh, pmmh, pdmmh, pmmh, ddmh, ddmh, and pmdm, respectively. DNA-binding dissociation rates are critphh, critpd, critpmh, and critpmh for Her-Her, Notch (NICD), Mespa-Mespa, and Mespb-Mespb, respectively. Transcriptional time delays of her, deltaC, mespa, and mespb mRNAs include the transcription, splicing, and nuclear-to-cytoplasmic transport, and these delays are represented by delayth, delaymd, delaymh, and delaymh, respectively. The translational time delays of Her, Mespa and Mespb proteins include translation and nuclear import of these repressor proteins, and these delays are represented by delayph, delayph, and delayph, respectively. The translational time delay of DeltaC protein includes translation and localization of DeltaC protein at the membrane, interaction of Delta-Notch proteins, and production and localization of NICD at the nucleus, and is expressed as delaypd. Likewise, we defined the time-delay δM, to represent the total time-delay in the indirect regulatory interactions between Mesp, Rippy1, and Tbx6 proteins (Windner et al., 2015). We represent the kth cell as c_k and time as t.

Delay Differential Equation Model:

A. mRNA Levels
A.1. her mRNA Levels

\[
\frac{\partial m_h(c_k, t)}{\partial t} = m_{sh} \left( \frac{1}{1 + \sum_{c_n \in N} \frac{pd(c_n, t - delay_{m_h})}{critpd}} \right) \left[ 1 + \sum_{c_n \in N} \frac{pd(c_n, t - delay_{m_h})}{critpd} + \left( \frac{phh(c_k, t - delay_{m_h})}{critphh} \right)^2 \right] - md_h \cdot mh(c_k, t)
\]

A.2. deltaC mRNA Levels

\[
\frac{\partial m_d(c_k, t)}{\partial t} = m_{sd} \left( \frac{1}{1 + \sum_{c_n \in N} \frac{pd(c_n, t - delay_{m_d})}{critphh}} \right) - md_d \cdot md(c_k, t)
\]

A.3. mespa mRNA Levels

\[
\frac{\partial m_{mm_a}(c_k, t)}{\partial t} = m_{sm_a} \left( \frac{1}{1 + \sum_{c_n \in N} \frac{pd(c_n, t - delay_{m_{mm_a}})}{critpd}} \right) \left[ 1 + \sum_{c_n \in N} \frac{pd(c_n, t - delay_{m_{mm_a}})}{critpd} + \left( \frac{phh(c_k, t - delay_{m_{mm_a}})}{critphh} \right)^2 \right] - md_{mm_a} \cdot m_a(c_k, t)
\]

A.4. mespb mRNA Levels

\[
\frac{\partial m_{mm_b}(c_k, t)}{\partial t} = m_{sm_b} \left( \frac{1}{1 + \sum_{c_n \in N} \frac{pd(c_n, t - delay_{m_{mm_b}})}{critpd}} \right) \left[ 1 + \sum_{c_n \in N} \frac{pd(c_n, t - delay_{m_{mm_b}})}{critpd} + \left( \frac{pm_{mm_a}(c_k, t - delay_{m_{mm_b}} - \delta_M)}{critpm_{mm_a}} \right)^2 \right] - md_{mm_b} \cdot m_b(c_k, t)
\]

where N represents all the neighbors of the kth cell (c_k)
B. Monomer Protein Levels

B.1. Her Monomer Protein Levels

\[ \frac{\partial p_h(c_k, t)}{\partial t} = psk \cdot mh(c_k, t - delay) + pdh \cdot ph(c_k, t) + 2dhh \cdot phh(c_k, t) - 2dahh \cdot ph(c_k, t) \cdot ph(c_k, t) \]

B.2. Delta Monomer Protein Levels

\[ \frac{\partial p_d(c_k, t)}{\partial t} = psd \cdot md(c_k, t - delay) - pdp \cdot pd(c_k, t) \]

B.3. Mespa Monomer Protein Levels

\[ \frac{\partial p_m_a(c_k, t)}{\partial t} = psm_a \cdot mm_a(c_k, t - delay) - pdm_a \cdot pm_a(c_k, t) + 2ddm_a \cdot pm_a(c_k, t) - 2dam_a \cdot pm_a(c_k, t) \cdot pm_a(c_k, t) + ddm_a_m_b \cdot pm_a(c_k, t) - dam_a_m_b \cdot pm_a(c_k, t) \cdot pm_b(c_k, t) \]

B.4. Mesph Monomer Protein Levels

\[ \frac{\partial p_m_b(c_k, t)}{\partial t} = psm_b \cdot mm_b(c_k, t - delay) - pdm_b \cdot pm_b(c_k, t) + 2ddm_b \cdot pm_b(c_k, t) - 2dam_b \cdot pm_b(c_k, t) \cdot pm_b(c_k, t) + ddm_a_m_b \cdot pm_a(c_k, t) - dam_a_m_b \cdot pm_a(c_k, t) \cdot pm_b(c_k, t) \]
C. Dimer Protein Levels

C.1. Her-Her Dimer Protein Levels

\[
\frac{\partial phh(c_k,t)}{\partial t} = -ddhh \cdot phh(c_k,t) + dahh \cdot ph(c_k,t) \cdot ph(c_k,t) - pdhh \cdot phh
\]

C.2. Mespa-Mespa Dimer Protein Levels

\[
\frac{\partial pm_a m_a(c_k,t)}{\partial t} = -ddm_a m_a \cdot pm_a m_a(c_k,t) + dam_a m_a \cdot pm_a(c_k,t) \cdot pm_a(c_k,t) - pdm_a m_a \cdot pm_a m_a
\]

C.3. Mesph-Mesph Dimer Protein Levels

\[
\frac{\partial pm_b m_b(c_k,t)}{\partial t} = -ddm_b m_b \cdot pm_b m_b(c_k,t) + dam_b m_b \cdot pm_b(c_k,t) \cdot pm_b(c_k,t) - pdm_b m_b \cdot pm_b m_b
\]

C.4. Mespa-Mesph Dimer Protein Levels

\[
\frac{\partial pm_a m_b(c_k,t)}{\partial t} = -ddm_a m_b \cdot pm_a m_b(c_k,t) + dam_a m_b \cdot pm_a(c_k,t) \cdot pm_b(c_k,t) - pdm_a m_b \cdot pm_a m_b
\]

Spatial modeling

A two-dimensional hexagonal grid of 4 x 50 cells was used to represent the PSM tissue in our simulations. The right- and left-most cells in each column were connected artificially, such that each cell has six neighbors, excluding the cells located in the most posterior and anterior columns that have only four neighbors. The model was simulated for 930 min in total. In the first 300 min, we simulated only 4 x 10 cells forming the posterior PSM. Then, we grew the posterior PSM tissue for 240 min until the PSM was full by adding a column of 4 cells every six minutes. After the PSM was full, we added a column of 4 cells at the posterior end, and removed an older column of cells at the anterior end to keep a fixed PSM size. We set the Her and DeltaC translational time delays within a biologically relevant range in the posterior PSM (first 10 columns of cells); this rate was linearly interpolated at all intermediate (middle 20 columns of cells) and anterior PSM locations (last 20 columns of cells). The largest translational time delays at the anterior PSM for Her and DeltaC proteins were set to 3.9 fold of posterior PSM (Ay et al., 2014). Similarly, the translational time delays of Mespa and Mesph proteins were increased 2.1 fold from mid-PSM to anterior PSM. The model implicitly implements the input of FGF signaling on mesp transcription, by restricting the transcription zone of mesp genes only to anterior PSM in simulations.

Pseudo-stochastic numerical simulation

We carried out pseudo-stochastic simulations of our delay differential equation model to reproduce randomness in the regulatory network. Biochemical reaction rates (parameters in the model) were allowed to change up to 16% between cells to generate the inherent stochasticity in this biological system. The variations in reaction rates were formed during the creation of the cells and kept constant throughout the lifetime of the cells in the PSM. The perturbed system of DDEs was solved numerically using Euler’s method. Euler’s method increments the time by the
chosen step size (0.01 min), and updates mRNA and mono and dimer protein levels after each iteration using the rates of change specified by the DDEs. To simulate a notch1a−/− mutant, we set the translation rate of the DeltaC protein (psd) to zero. To simulate the DAPT condition, we set psd to 0 after 600 min. Overexpression of her gene was modeled by increasing its translation rate psh from 600 to 630 min. Similarly, we modeled overexpression of mespa and mespb genes by increasing their translation rates psma and psmb from 600 to 660 min.

Parameter estimation

Delays in transcription and translation and degradation rates of mRNA and protein have been measured experimentally (Ay et al., 2014; Ay et al., 2013; Giudicelli et al., 2007; Hanisch et al., 2013). However, some of the reaction rates have not been measured due to technical limitations. We used parameter estimation to identify biologically relevant reaction rates (model parameters) that could reproduce the experimental observations.

Parameter search was performed using the stochastic ranking evolutionary strategy (SRES) algorithm (Runarsson and Yao, 2000), which looked for suitable parameter sets fit to experimental conditions (Table S2). The SRES algorithm performs better than other parameter estimation algorithms in large-scale non-linear biological systems (Fakhouri et al., 2010; Fomekong-Nanfack et al., 2007; Moles et al., 2003). We used the ranges provided in Ay et al. (2014) for the parameters that represent the same functionality as the model from (Ay et al., 2014). New parameters were given freedom within biologically realistic limits. To produce biologically feasible parameter ranges (Table S1), parameter ranges were refined a few times based on the results of initial parameter searches.

Posterior PSM

The DDE model was simulated for 300 min in 40 (4 rows of 10 cells) posterior PSM cells to obtain her mRNA expression levels. The period was calculated as the time difference between the last two peaks of her mRNA oscillations.

Whole PSM

The PSM reached its full size of 200 cells in 540 min in our simulations. To calculate the amplitude and synchrony of segmentation network genes, we took 1 snapshot for overexpression experiments, and 5 snapshots for notch1a−/− mutant and DAPT treatment over 30 min of simulation.

The amplitudes for overexpression embryos were calculated as the change between the average of top ten and bottom ten corresponding gene expression levels. The amplitudes for notch1a−/− mutant and DAPT embryos were calculated as the mean of the five measured amplitudes found using five snapshots. The synchronization scores for overexpression transgenics were measured by finding average of the three Pearson correlation coefficients measured between each row of cells and the first row of cells. The synchronization scores for wildtype, notch1a−/−, and DAPT embryos were calculated as the mean of the five calculated synchronization scores found using five snapshots. The complementarity between her and mespa/mespb genes were calculated as the average of the fifteen (five snapshots, four rows) Pearson correlation coefficients between her and mespa/mespb gene expression levels.

Coding

The model and related analysis are implemented in C++ and Python. C++ is used for the model’s numerical simulation and parameter search, because of its speed. Python is used due to its user-friendly and superior data processing and plotting libraries. Our code can perform a 930
min simulation of 200 PSM cells in less than 1 min on an iMAC running MAC OS X 10.12.4 with 3.1 GHz Intel Core i7 and 16 GB of RAM. A parallel version of our code is written using the Message Passing Interface (MPI) for time-intensive parameter searches. Each SRES parameter estimation run with a population size of 20, 3 parents, and 2,000 generations took approximately 48 hours using 24 processors on a computer cluster of 19 nodes, 248 processors, and 24 gigabytes of RAM per node.

DATA AND SOFTWARE AVAILABILITY
The C++ and Python codes will be made available upon request.

REFERENCES
Ay, A., Holland, J., Sperlea, A., Devakanmalai, G.S., Knierer, S., Sangervasi, S., Stevenson, A., and Ozbudak, E.M. (2014). Spatial gradients of protein-level time delays set the pace of the traveling segmentation clock waves. Development 141, 4158-4167.
Ay, A., Knierer, S., Sperlea, A., Holland, J., and Özbudak, E.M. (2013). Short-lived Her Proteins Drive Robust Synchronized Oscillations in the Zebrafish Segmentation Clock. Development 140, 3244-3253.
Fakhouri, W.D., Ay, A., Sayal, R., Dresch, J., Dayringer, E., and Arnosti, D.N. (2010). Deciphering a transcriptional regulatory code: modeling short-range repression in the Drosophila embryo. Molecular Systems Biology 6.
Fomekong-Nanfack, Y., Kaandorp, J.A., and Blom, J. (2007). Efficient parameter estimation for spatio-temporal models of pattern formation: case study of Drosophila melanogaster. Bioinformatics 23, 3356-3363.
Giudicelli, F., Ozbudak, E.M., Wright, G.J., and Lewis, J. (2007). Setting the Tempo in Development: An Investigation of the Zebrafish Somite Clock Mechanism. PLoS Biol 5, e150.
Hanisch, A., Holder, M.V., Choorapoikayil, S., Gajewski, M., Ozbudak, E.M., and Lewis, J. (2013). The elongation rate of RNA Polymerase II in the zebrafish and its significance in the somite segmentation clock. Development 140, 444-453.
Henry, C.A., Urban, M.K., Dill, K.K., Merlie, J.P., Page, M.F., Kimmel, C.B., and Amacher, S.L. (2002). Two linked hairy/Enhancer of split-related zebrafish genes, her1 and her7, function together to refine alternating somite boundaries. Development 129, 3693-3704.
Jiang, Y.J., Aerne, B.L., Smithers, L., Haddon, C., Ish-Horowicz, D., and Lewis, J. (2000). Notch signalling and the synchronization of the somite segmentation clock. Nature 408, 475-479.
Lee, Y., Grill, S., Sanchez, A., Murphy-Ryan, M., and Poss, K.D. (2005). Fgf signaling instructs position-dependent growth rate during zebrafish fin regeneration. Development 132, 5173-5183.
Lewis, J. (2003). Autoinhibition with transcriptional delay: A simple mechanism for the zebrafish somitogenesis oscillator. Current Biology 13, 1398-1408.
Lewis, J.L., Bonner, J., Modrell, M., Ragland, J.W., Moon, R.T., Dorsky, R.I., and Raible, D.W. (2004). Reiterated Wnt signaling during zebrafish neural crest development. Development 131, 1299-1308.
Moles, C.G., Mendes, P., and Banga, J.R. (2003). Parameter estimation in biochemical pathways: A comparison of global optimization methods. Genome Research 13, 2467-2474.
Oates, A.C., and Ho, R.K. (2002). Hairy/E(spl)-related (Her) genes are central components of the segmentation oscillator and display redundancy with the Delta/Notch signaling pathway in the formation of anterior segmental boundaries in the zebrafish. Development 129, 2929-2946.
Ozbudak, E.M., and Lewis, J. (2008). Notch signalling synchronizes the zebrafish segmentation clock but is not needed to create somite boundaries. PLoS genetics 4, e15.
Riedel-Kruse, I.H., Muller, C., and Oates, A.C. (2007). Synchrony dynamics during initiation, failure, and rescue of the segmentation clock. Science 317, 1911-1915.
Runarsson, T.P., and Yao, X. (2000). Stochastic ranking for constrained evolutionary optimization. Ieee T Evolut Comput 4, 284-294.
Takke, C., and Campos-Ortega, J.A. (1999). her1, a zebrafish pair-rule like gene, acts downstream of notch signalling to control somite development. Development 126, 3005-3014.
Watanabe, T., Saito, D., Tanabe, K., Suetsugu, R., Nakaya, Y., Nakagawa, S., and Takahashi, Y. (2007). Tet-on inducible system combined with in ovo electroporation dissects multiple roles of genes in somitogenesis of chicken embryos. Dev Biol 305, 625-636.
Wehner, D., Jahn, C., and Weidinger, G. (2015). Use of the TetON System to Study Molecular Mechanisms of Zebrafish Regeneration. Jove-J Vis Exp.
Windner, S.E., Doris, R.A., Ferguson, C.M., Nelson, A.C., Valentin, G., Tan, H., Oates, A.C., Wardle, F.C., and Devoto, S.H. (2015). Tbx6, Mesp-b and Ripply1 regulate the onset of skeletal myogenesis in zebrafish. Development 142, 1159-1168.