Presomitic mesoderm-specific expression of the transcriptional repressor Hes7 is controlled by E-box, T-box, and Notch signaling pathways

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Establishment of cellular identities is essential for generating complex patterns during embryogenesis. To establish cellular identities and develop organ/tissue formations properly, gene expressions are spatiotemporally regulated with accuracy in appropriate domains. The lefty1/lefty2 expressions, for example, are restricted to the left side of early developing mouse embryos to direct left–right axis determination (1). Deletion of lefty1 or lefty2 results in left pulmonary isomerism, malpositioning of the cardiac outflow tracts, and other vascular vessels (2) or an expanded primitive streak, formation of excess mesoderm (3), and various situs defects, including left isomerism (4), respectively.

We have identified the Hes7 gene, one of the Hes family transcriptional repressors, which is exclusively expressed in the presomitic mesoderm (PSM) and acts as a key molecule for somitogenesis (5–7). Somitogenesis is the process to form somites, which is a pair of epithelial spheres beside a neural tube and appear transiently during embryogenesis, from the anterior PSM (8). It is known that Hes7 expression is restricted in the PSM and is regulated by the Notch, Fgf, and Wnt signaling pathways (9). These signaling pathways regulate various processes during embryogenesis, suggesting that the restricted Hes7 gene expression in the PSM is orchestrated by a combination of transcriptional factors downstream of the Notch, Fgf, and Wnt signaling pathways. However, little is known about the transcriptional regulations that are associated with Hes7 gene expression.

In this study, we describe the presence of a Hes7 essential region, residing from −1.5 to −1.1 kb from the transcription start site of the mouse Hes7 gene, that directs PSM-specific Hes7 expression. Furthermore, we demonstrate the mechanisms for Hes7 expression in the PSM. Restricted Hes7 expression is controlled through E-box, T-box, and the RBPj-binding element in the Hes7 essential region, presumably activated by a synergistic effect of mesogenin1, Tbx6, and Notch signaling, and repressed by Tbx18, Ripply2, and Hes7. Our study uncovered that the Hes7 essential region directs PSM-restricted expression pattern of Hes7, orchestrated by multiple transcriptional elements.

Results

C region, from −1.5 to −1.1 kb upstream of TSS of mouse Hes7, is sufficient for accurate Hes7 expression in the PSM

Although Hes7 mRNA is well known to be exclusively expressed in the PSM, the molecular mechanisms that regulate Hes7 expression/repression remain largely unknown. To uncover molecular mechanisms of the unique Hes7 expression in the PSM, we first tried to search for the essential region for

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This article contains Figs. S1–S10 and supporting Experimental procedures.
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The abbreviations used are: PSM, presomitic mesoderm; TSS, transcription start site; FW, forward; RV, reverse; bHLH, basic helix-loop-helix; PVDF, polyvinylidene difluoride.
PSM-specific Hes7 expression by means of transgenic founder assays. All transgenic embryos \((n = 3)\) carrying the 5.3-kb fragment upstream from the TSS of mouse Hes7, which has been utilized for exogenous Hes7 expression in the PSM (10, 11), followed by a lacZ reporter showed X-gal–positive staining in the PSM at embryonic day (E) 10.5 (Fig. 1B). We next narrowed down the essential region and found that 2.4- and 1.5-kb fragments upstream from TSS were still sufficient for the reporter expression in the PSM (Fig. 1, C and D). However, we could not detect any X-gal–positive staining in the PSM of transgenic mice carrying a 1.1-kb fragment or shorter fragment (Fig. 1, E and F). These results suggest that the 1.5-kb fragment is sufficient for the PSM-specific expression of Hes7 and that the essential region for PSM-restricted expression resides between \(-1.5\) and \(-1.1\) kb upstream of TSS.

To confirm the region responsible for PSM-specific expression of Hes7, we next investigated the fragment from \(-2.4\) to \(-1.5\) kb, which is not considered to be an essential region for PSM-specific Hes7 expression, and hereafter it is referred to as the B region; and the fragment from \(-1.5\) to \(-1.1\) kb is hereafter referred to as the C region (Fig. 1A). Transgenic founder assays revealed that the B region drove X-gal protein expression in the PSM, whereas the B region had no \(\beta\)-gal activity in the PSM (Fig. 1, G and H). Next, we narrowed down the essential
region in the C region. We constructed four reporter vectors deleting a quarter of fragment C region named Deletion1, -2, -3, or -4 (Fig. 1A and Fig. S1). Deletion1 and -2 showed no X-gal-positive staining at the most posterior end of the PSM (Fig. 1, I–L). However, Deletion3 and -4 resulted in X-gal-positive staining comparable with the WT C region (Fig. 1, M–P). In situ hybridization also demonstrated that the anterior-most regions of the PSM were negative for Deletion1- and -2-driven lacZ mRNA (Fig. S2). We therefore deduce from the above results that the 0.4-kb C region is the essential region, and the distal half of the C region from TSS contains essential transcriptional binding sites for Hes7 expression in the PSM.

Although endogenous Hes7 mRNA was expressed exclusively in the PSM, X-gal-positive staining was present throughout the PSM and newly formed somites, which are derived from the PSM, in transgenic mice carrying 5.3-, 2.4-, 1.5-kb fragments or the C region (Fig. 1, B–D and H). A simple explanation was that the reporter mRNA was transcribed exclusively in PSM cells, whereas its resultant β-gal protein remained in differentiated somite cells due to high protein stability. To assess this possibility, we carried out lacZ mRNA detection by in situ hybridization. The lacZ mRNA driven by a 5.3-kb fragment or the C region was exclusively expressed in the PSM (Fig. 1, R and T), and as expected, the control mice without transgene did not show any signal in the PSM (Fig. 1, Q and S). These results demonstrate that the C region is sufficient for accurate Hes7 expression in the PSM.

E-boxes in C region are essential to drive Hes7 in the PSM

To address which transcriptional binding elements regulate PSM-specific Hes7 expression, we searched for putative transcriptional binding elements within the C region by in silico analyses. There are several putative regulatory sequences in the C region: three T-boxes (YMACCYY or complementary) (21); six E-boxes (CANNTG) (5); and one RBPj-binding site (YRTGDGAD or complementary) (24). In particular, E-box1, -3, -5, and -6 and T-box2 were completely conserved among Homo sapiens, Pongo abelii, Bos taurus, and Mus musculus (Fig. 2A). To address whether these putative E-boxes, T-boxes, and the RBPj-binding site in the C region are functional in vivo, we made transgenic mice carrying the mutated E-box1–6/C region or the mutated T-box1 and -2/C region, which also include a mutated RBPj-binding site. In situ hybridization assays using transgenic founder mice revealed that mice with the mutated T-box1 and -2/C region expressed the reporter mRNA in the PSM as well as the mice with WT C region (Fig. 2B, panels a and b). In contrast, 14 of 16 mice with the mutated E-box1–6/C region showed no positive signal, whereas only two embryos showed a dispersed and straggling reporter expression (Fig. 2B, panels c and d, and Fig. S3), indicating that E-boxes in the C region are essential to drive Hes7 in the PSM.

Msgn1, Tbx6, and Notch signaling pathway activate C region in vitro

To investigate the putative elements in the C region in depth, we performed luciferase assays using constructs containing the WT or mutated C region followed by a human β-globin minimal promoter. To perform this, mesogenin1 (Msgn1), Tbx6, and NICD, the intracellular domain of Notch1, were utilized as binding factors for E-box, T-box, and RBPj-binding site, respectively. This is based on previous reports that showed that Msgn1, a bHLH-type transcription factor, is exclusively expressed in the posterior PSM (12), whereas Tbx6, a T-box family of transcriptional factor, and Notch signaling molecules are expressed ubiquitously in mouse PSM (13–15). We confirmed that the Msgn1 expression domain coincided with Hes7 stripes in phase I and II, but not at phase III, whereas Tbx6 mRNA was constantly distributed over the PSM, overlying any phases of the Hes7-transcribed region, besides the anterior-most region of the PSM (Fig. S4). WT C region reporter was activated by all of the transcriptional factors tested (Fig. 3, A–C), whereas the activity of mutated E-box1–6/C or T-box1–3/C region reporter was attenuated by Msgn1 or NICD, respectively. (Fig. 3, A and B), indicating that at least one of the mutated E-boxes is a functional site for Msgn1, whereas the T-box1 is receptive toward Notch signaling molecules. However, the mutated T-box1–3/C region reporter was still responsive to Tbx6, comparable with that of WT C region (Fig. 3C).

To further investigate E-boxes, T-boxes, and the RBPj-binding site in the C region in vitro, we performed oligo-DNA pulldown assay and electrophoretic mobility shift assay (EMSA). Pulldown assay revealed that Msgn1 bound to E-box1, but not to E-box2 and -3 (Fig. 3D). Furthermore, we confirmed that Msgn1 binding to the E-box1 was abolished by the mutation of E-box1 (Fig. 3D). EMSA also demonstrated that Msgn1 bound to E-box1, but not to E-box2 and -3 (Fig. S5). Again, we confirmed that Msgn1 binding to E-box1 was abolished by an excess amount of the nonlabeled E-box1 but not by that of the mutant one (Fig. S5). These results raise the possibility that Msgn1 in the PSM activates Hes7 expression via the E-box1 in C region. RBPj binding was detected by T-box1, in which RBPj-binding site is included, by pulldown assay and EMSA (Fig. 3E and Fig. S6). Moreover, pulldown assay demonstrated that RBPj binding was dramatically attenuated by the mutated E-box1, and EMSA showed that this binding was attenuated by an excess amount of the nonlabeled E-box1 but not by that of the mutated one (Fig. 3F and Fig. S6). These results raise the possibility that the RBPj-binding site in C region is functional in vivo. Luciferase assays demonstrated that T-box1–3 were not responsive to Tbx6; however, ChIP assay utilizing PSM samples indicated that Tbx6 bound to the C region (Fig. S7A). To investigate whether Tbx6 binds to T-box elements in the C region, we performed oligo-DNA pulldown assay and showed that Tbx6 bound to T-box1 and T-box2 but not to T-box3. In contrast, the Tbx6 binding potential to T-box1 or T-box 2 was eliminated by mutated T-box1 or T-box2, respectively (Fig. S7 and data not shown).

Msgn1, Tbx6, and Notch signaling pathways synergistically activate C region in vitro

As we demonstrated that Msgn1 and NICD increased the luciferase activity of the C region and Msgn1 and RBPj bound to E-box1 and T-box1, respectively (Fig. 3, A–F), we next investigated whether a combination of Msgn1 and NICD show a coordinated activation. Compared with single Msgn1 or NICD
activation, co-expression of Msgn1 and NICD synergistically increased luciferase activity (Fig. 3F). Although, as shown above, T-box1 and -2 were not necessary for the expression of Hes7 in the PSM (Fig. 2B, panels a and b), the ChiP assay, oligo-DNA pulldown assay, and luciferase assay showed that Tbx6 could bind to and activate the C region (Fig. 3C and Fig. S7). Furthermore, because Tbx6 has been known to work synergistically with other transcriptional activators for gene expressions in the PSM (13), we investigated the possibility of Tbx6 working synergistically with Msgn1 or NICD to activate the C region. In contrast to either Msgn1 or NICD alone, the combination of Tbx6 with either Msgn1 or NICD increased the C region-driven luciferase activity (Fig. 3G). Intriguingly, we revealed that Tbx6 together with Msgn1 and NICD accelerated the reporter expression much more than the combinations with Tbx6/Msgn1, Tbx6/NICD, or Msgn1/NICD (Fig. 3G). These results indicate that Tbx6, Msgn1, and Notch signaling activate Hes7 expression coordinately via the C region at least in vitro. To confirm whether their synergistic effect on the C region is due to E-boxes, T-boxes, and the RBPj-binding site, we next performed luciferase assays using the C region with mutated T-box1–3 or E-box1–6/C region. Synergistic activation by Msgn1/NICD/Tbx6 was almost completely abolished in T-box or E-box mutants (Fig. 3H), although activity in the T-box1–3 mutant was higher than that of the E-box1–6 mutant, supporting the results that embryos with mutated
Hes7 expression, although T-box1–3 elements were not necessary for the single activation by Tbx6 (Fig. 3C). In contrast, the activation of the C region with E-box1–6 mutant by Tbx6 and Msgn1 was completely abolished (Fig. S8), which was similar to the result by Msgn1/NICD/Tbx6 (Fig. 3H), indicating that E-boxes are essential for the synergistic Hes7 expression. Taking these in vitro and in vivo results together (Figs. 2 and 3), we deduce that E-boxes and T-boxes, including the RBPj-binding site, are critical and auxiliary for Hes7 expression, respectively.

Tbx18, Ripply2, and Hes7 repress the activation of C region in vitro

Transgenic mice carrying the mutated T-box1 and -2/C region or Deletion1 expressed lacZ mRNA not only in the PSM but also expressed a dispersed and straggling reporter mRNA in the somites (Fig. 2B, panel b, and Fig. S2), suggesting that the C region, especially T-boxes, has a role in preventing ectopic Hes7 expression. To understand the molecular mechanisms of how Hes7 expression in the anterior-most PSM or in somites is suppressed, we next investigated the repression mechanisms for Hes7. Tbx18 is one of the transcriptional repressors among the T-box family of transcriptional factor genes that is expressed in mouse PSM (13), and as reported previously (16), Tbx18 is expressed in the rostral part of somites and the anterior-most PSM where Hes7 propagation has vanished (Fig. S4, C and C’), demonstrating that Tbx18 and Hes7 expressions are mutually exclusive. We investigated whether Tbx18 binds to T-boxes in the C region by oligo-DNA pulldown assay and EMSA. The pulldown assays demonstrated that Tbx18 bound to WT T-box1 but not to T-box2 or -3 (Fig. 4A and Fig. S9A). Tbx18 binding to T-box1 was diminished by the mutation of T-box1 (Fig. 4A). EMSA also showed that Tbx18 bound to T-box1 but not to T-box2 or -3 (Fig. S9B). Again, we demonstrated that Tbx18 binding to T-box1 was weakened by an excess amount of the nonlabeled T-box1 but not by that of the mutated one (Fig. S9B). Furthermore, luciferase assays revealed that Tbx18 dose-dependently repressed reporter activity induced by Tbx6 and NICD (Fig. 4B), which are expressed at the anterior-most PSM (Fig. 5 and Fig. S4).

Ripply2 mRNA was strongly expressed, as reported previously (17), in the anterior PSM (prospective somites S0 and S-I) when Hes7 was in phase I (Fig. S4D), whereas in phase III of Hes7, Ripply2 showed two weak stripes at the region where Hes7 is lost (Fig. S4D’). Because Ripply co-repressors have been known to act on the repressor by interacting with Xenopus Tbx6 or zebrafish Tbx24, which are structurally related to mouse Tbx6 (18, 19), we performed co-immunoprecipitation assays in culture cells, HEK293T, to examine whether mouse Ripply co-repressors interact with mouse Tbx6. We revealed that Ripply1/2 form a complex with Tbx6 in vitro (Fig. 4C), especially Ripply2, which had a high affinity to Tbx6. Luciferase assay further uncovered that luciferase activities of the C region induced by Tbx6/NICD were reduced by Ripply2 (Fig. 4B). Moreover, co-expression of Tbx18 and Ripply2 repressed luciferase activity more effectively compared to when either Tbx18 or Ripply2 was expressed, suggesting that Tbx18 and Ripply2 repress the C region independently. These results raise the pos-
sibility that Tbx18 and Ripply2 are repressive regulators for Hes7 termination in the anterior-most PSM and somites by binding to T-box1 or by forming a complex with Tbx6 to reduce Tbx6 transcriptional activity, respectively.

Because Hes7 could bind to the E-box to repress transcriptional activity, we investigated whether E-boxes in the C region are functional for Hes7. We first demonstrated that Hes7 binding to the C region in vivo was detected by ChIP assay using mouse PSM (Fig. 4D). Furthermore, the oligo-DNA pulldown assay showed that Hes7 could bind to E-box2 and -3 but not to E-box1 (Fig. 4E). Moreover, luciferase assays demonstrated that Hes7 repressed synergistic Msgn1/Tbx6/NICD activation dose-dependently (Fig. 4F). These results suggest that the C region might also be associated with the oscillatory expression of Hes7.

Finally, we sought to investigate whether epigenetic regulations, especially DNA methylation, of the C region take part in the repression of Hes7 ectopic expression. To that end, we examined the DNA methylation status of the C region by bisulfite sequencing. However, no CpG sites within the C region were methylated in all the tissues tested, including PSM, head, and caudal trunk from E10.5 embryo (Fig. S10). Experimental procedures for this bisulfite sequence are provided in the supporting Experimental procedures. This result suggests that the regulation by DNA methylation is not correlated with the mechanism for the PSM-specific expression of Hes7.

Discussion

Spatiotemporal Hes7 expression pattern is very unique, whereby it is restricted to the PSM and propagates (oscillates) from the posterior-end to the anterior-end of the PSM with 2-h periodicity in mice. In this study, we identified a narrow region within 1.5 to 1.1 kb from TSS in mouse Hes7, referred to as the C region in this study, that directs the specific expression of the reporter gene in the PSM during mouse embryogenesis. At the molecular level, we found E-boxes, T-boxes, and RBPj-binding sites in the C region and further demonstrated that these elements are crucial for the restricted expression of the reporter gene in the PSM. Furthermore, this study raises the possibility
that the Notch signaling pathway, the transcriptional activators, Msgn1 and Tbx6, and the transcriptional repressors, Tbx18, Ripply1/2, and Hes7, participate as novel factors for the C region's activation/repression.

Our study using transgenic founder assays revealed that this C region, which is highly conserved among several species, is sufficient to direct expression of the reporter gene in the PSM specifically during mouse embryogenesis (Figs 1 and 2). Our current study further dissected the molecular mechanisms for PSM-specific Hes7 expression. In situ hybridization for lacZ mRNA (Fig. 2B) showed that only two of 16 embryos with mutated E-box demonstrated positive, but obscure, signals. In addition, luciferase assays (Fig. 3H) demonstrated that the mutated E-box was completely unresponsive to the activation by Tbx6/Msn1/NICD. These findings suggest that the mutated E-box has no potential to activate gene expression in the PSM. In contrast to the mutated E-box, the mutated T-box had produced lacZ signals in six of eight embryos (Fig. 2B) and had increased the luciferase activity by Tbx6/Msn1/NICD (Fig. 3H), although the activity was much lower than that of WT C region. We therefore conclude that E-boxes and T-boxes, including the RBPj-binding site, are critical and auxiliary, respectively. Moreover, X-gal staining and in situ hybridization for Deletion1 or -2 (Fig. 1, 1–L, and Fig. S2) suggest that E-box1–3 and T-box1/2 are functional, because these results showed no/weak signals at the posterior-end PSM. The reason no/weak signals were restricted at the posterior-end PSM could be that the gene expression driven by the C region without any of these elements is very weak and under detectable levels by X-gal staining and in situ hybridization at the posterior-end PSM; however, after a while, reporter mRNA and protein slowly accumulated to reach detectable levels.

It has been reported that the somite formation does not take place when Msgn1 or Tbx6 is knocked out in mice (12, 20, 21). In addition to Msgn1 and Tbx6, somite formation does not occur properly when Hes7 is deficient (6). Moreover, Msgn1 or components of the Notch pathway knockout mice express less Hes7 in the PSM (10, 22, 23). These reports strongly support our findings that Msgn1, Tbx6, and Notch pathway are upstream of Hes7. Furthermore, we raise the possibility that the combination of Msgn1, Notch signaling, and Tbx6 induces Hes7 expression via the C region in the mouse PSM. Interestingly, synergistic transcriptional activation by Tbx6 with other transcriptional factors during somitogenesis has been reported; for example, Tbx6 cooperates with Wnt signaling for Dll1 and Msgn1 induction (24, 25) and with Notch signaling for Mesp2 induction in mouse (26). In Xenopus, Tbx6 activates bowlina, a Xenopus Ripply homologue, in synergy with bHLH transcription factors, Thylacine1 and E47 (27). Moreover, we also raise the possibility that T-box1–3 elements are essential to form the optimal three-dimensional structure of the C region with Tbx6/Msn1/NICD for the synergistic activation. Although further investigations will be required to address whether and how Tbx6/Msn1/NICD form a complex with the C region for the complete Hes7 activation, our finding nonetheless had shed light on a new role of Tbx6 in somitogenesis.

Our findings in this study also raised the possibility that T-box elements function to repress Hes7 expression in the anterior-most PSM (S-I and S0) and somites by binding with Tbx18. This dual function of T-boxes could be one of the mechanisms for the termination and inhibition of Hes7 expression at the anterior-most PSM and somites, respectively. Another key factor for Hes7 repression is Ripply1/2. Ripply1/2 have already been shown to repress Tbx6 expression by two different ways. One is the conversion of Xenopus and zebrafish Tbx6 from transcriptional activator to suppressor by binding with Groucho/TLE co-repressors (18, 19). The other is the elimination of Tbx6 protein by unknown mechanisms (28). In this study, we demonstrated that mouse Ripply1/2 could bind with mouse Tbx6 and repress the expression of Hes7 in vitro (Fig. 4, B and C). Our data suggest that, at the anterior-most PSM in mouse, Ripply suppresses Hes7 expression through recruitment to T-box elements with Tbx6/Groucho/TLE co-repressors and/or by eliminating Tbx6 protein (Fig. 5). However, because Hes7 expression patterns are normal even in Ripply1 and -2 double knockout embryos (28), the Hes7 termination mechanism by Ripply1/2 might be ancillary and that by Tbx18 is primary. It is noteworthy that an interesting paper has reported that Mesp2, which expresses S-I, suppresses Notch signaling via destabilizing Mastermind-like 1, a coactivator of Notch signaling (29). The suppression of Notch signaling by Mesp2 at the anterior-most PSM shown in that report could be another potential mechanism for the termination of Hes7 expression. Taking their findings and our current analyses together, we establish that to spatiotemporally express and terminate Hes7 expression at the PSM, a web of transcriptional mechanisms is required during somitogenesis and that the Hes7 suppressor element would be important for a proper maintenance of the functional Hes7 domain as well as activator elements.

González et al. (30) reported that Hes7 expression is controlled by Tbx6 and Wnt signaling. They have identified an essential 400-bp region (~1.4 to ~1.0 kbp from TTS) for proper Hes7 expression, which is almost identical to our Hes7 essential region (~1.5 to ~1.1 kbp from TSS). Furthermore, they have also found that the activity of the Hes7 promoter in mouse PSM requires Tbx6-binding sites within this 400-bp region. These finding support our current study, although their distal T-box corresponds to T-box2 in our study, and the proximal one, which we missed as T-box, overlaps with E-box3 identified in our study. Intriguingly, they have mentioned that downstream molecules of the Wnt pathway activate the Hes7 promoter cooperatively with Tbx6 in cell culture and are necessary for its proper expression in the mouse PSM. More interestingly, they have shown that the expression of Msgn1, one of the Wnt target genes and the activator for the Hes7 essential region revealed in our study, is activated in embryos treated with LiCl, the inhibitor of GSK3β, in which Wnt signaling is activated. Taken together, their study strongly supports our results that Msgn1 is associated with the activation of Hes7 expression with Tbx6.

Our current findings demonstrated that the C region, including the specific elements of the C region, is sufficient for the restricted reporter mRNA expression in the PSM, and the C region is activated by a synergistic effect of Msgn1, Tbx6, and Notch signaling and repressed by Tbx18, Ripply2, and Hes7 in cell culture. However, we have not addressed whether the C
Promoter elements for restricted expression of Hes7

region and the regulatory factors are indispensable for the endogenous Hes7 expression in the PSM. In addition, we cannot rule out the possibility that other existing shadow or cryptic enhancers such as long range (1 Mb or more) enhancer (31) and enhancer residing in introns (32) or in 3’ downstream (33) may also be essential to Hes7 expression. To uncover whether the C region, and which elements in the C region, are indispensable for the PSM-specific endogenous Hes7 expression, further in vivo transciptional analyses of the endogenous Hes7 promoter deleting the whole C region or knocking-in the C region with mutated elements will be required. As mentioned above, previous reports suggest that Msn1 and Notch pathway are upstream of Hes7; however, detailed analyses of endogenous Hes7 expression in gene-modified mice in which the expression levels of Msn1, Tbx6, components of Notch signaling, Tbx18, or Ripply2 are altered will be required.

Finally, although we demonstrated that (i) C region has multiple E-boxes, (ii) the bHLH-type activator Msn1 and the repressor Hes7 occupy E-box1 and E-box2 and -3, respectively (Figs. 3D and 4E), and (iii) Hes7 can repress Msn1/Tbx6/NICD-dependent activation (Fig. 4D). Although we were unable to demonstrate the oscillation by the 5.3-kb fragment (Fig. 1R), we were unable to evince that the C region is enough for Hes7 oscillation, in that we could not detect the oscillatory lacZ mRNA expression pattern driven by the C region (Fig. 17). As it has been known that multiple enhancers act on gene expression to ensure robustness (34), multiple additional elements, such as E/N-boxes near the transcription start site as reported previously (35), might be needed for Hes7 to achieve and maintain oscillatory expression. Further analysis therefore will be required to find a minimum set of oscillatory elements for establishment of Hes7 oscillatory expression.

Experimental procedures

Animals

CD1 mice used in this study were purchased from SLC (Japan). Our experiments with mice have been approved by The Animal Care Committee of Nara Institute of Science and Technology (NAIST). These experiments were conducted in accordance with guidelines that were established by the Science Council of Japan.

Reporter constructs and transgenic mice

Hes7 upstream region was cloned by conventional molecular biological methods. Upstream fragment was PCR-amplified and inserted into pBluescriptII (Stratagene) with lacZ gene and SV40 poly(A) signal. Human β-globin minimal promoter, being synthesized as a double-stranded oligonucleotide, was inserted into a reporter vector for the enhancer assay. Sequences for human β-globin minimal promoter were as follows: Fw, TCC CGG CCT GGG CAT AAA AGT CAG GGC AGA GCC ATC TAT TGC TTA CAT TTG CTT C, and Rv, GAA GCA AAT GTA AGC AAT AGA TGG CTC TGC CCT GAC TTT TAT GCC CAG CCC GGG A. To make transgenic mice, the constructs were linearized and injected into fertilized eggs from CD1 mice by the animal facility of NAIST.

Luciferase assay

Hes7 C region followed by human β-globin minimal promoter were inserted into pGL3-Basic vector (Promega). Transcription factors were cloned by PCR with cDNA from mouse PSM. 5’-UTR and the coding region of each gene were inserted into pcDNA3 (Invitrogen), including FLAG, HA, or Myc tag at the 3’-end. 3 × 10⁴ NIH3T3 cells were plated in each well of a 24-well plate and were cultured in 10% fetal bovine serum/Dulbecco’s modified Eagle’s medium at 5% CO₂. After 24 h, cells were co-transfected with 300 ng of Hes7 reporter and 200 ng of expression vector of transcription factors using TransIT LT1 (Mirus). Transfected cells were lysed after 24 h of culture, and reporter activity was measured using a Dual-Luciferase assay system (Promega) and analyzed by ARBO (PerkinElmer Life Sciences). Firefly luciferase activity of the reporter was normalized by the activity of Renilla luciferase under control of SV40 promoter.

Mutagenesis

Site-directed mutagenesis was performed as described previously (36). Sequences were substituted as follows: T-box1, AGG TGT GGG AA to AGG TtT taa Ac; T-box2, CCA CAC CC to CgA tAt CC; T-box3, CCA CAC AT to CgA tAt AT; E-box1, CAT ATG to gtT Aac; E-box2, CAC GTG to gtC Gac; E-box3, CAG GTG to gtG Gac; E-box4, CAT CTG to gtT Cac; E-box5, CAT TTG to gtT Tac; and E-box6, CAA ATG to gtA Aac. Small letters indicate mutated nucleotides.

In silico promoter analysis

A homology between human Hes7 upstream and mouse Hes7 upstream was analyzed by VISTA (37, 38). To predict T-boxes, E-boxes, or RBpJ-binding sites in the C region, YMACCRY or complementary, CANNTG, or YRTGDAG or complementary were referred to as T-box, E-box, or RBpJ-binding site, respectively (5, 21, 39). Homology search among C regions of H. sapiens, P. abelii, B. taurus, and M. musculus was performed by ClustalW version 2.1.

Oligo-DNA pulldown assay

COS7 cells were seeded in 10-cm dish (4 × 10⁵ cells/dish), transfected with 15 μg of expression vectors, and cultured for 48 h. Cells were lysed in binding buffer (10 mm Tris, pH 8.0, 150 mm NaCl, 1 mm MgCl₂, 0.5% Nonidet P-40, 5% glycerol) with protease inhibitor mixture (Nacalai Tesque, Japan). After removing debris by centrifugation, 30 μl of 50% slurry streptavidin-Sepharose beads (GE Healthcare) and 200 pmol of dou-
ble-stranded oligonucleotide conjugated with biotin at each 5'-end were added to the cell lysates. The mixture was incubated at 4 °C for 30 min with rotation. Sepharose beads were washed with the binding buffer three times. Then, the resultant pulldown samples were separated by SDS-PAGE on precast 5–20% polyacrylamide gels (Nacalai Tesque, Japan). Proteins were then transferred to PVDF membranes using a wet electroblotting apparatus. The membranes were blocked using 5% skim milk in TBS with 0.1% Tween (TBS-T) for 1 h and incubated with anti-Myc (monoclonal, PL14, MBL, Japan) overnight, followed by incubation with anti-mouse IgG conjugated with horseradish peroxidase (GE Healthcare). Signals were visualized by the enhanced chemiluminescence detection system according to the manufacturer's instruction (Nacalai Tesque, Japan). Oligo-DNA sequences are as follows: E-box1 FW, 5'-AAA GTC ATT CCA TAT GGC CAG GGG CG-3’, and E-box1 RV, 5’-CGC CCC TGG CCA TAT GGA ATG ACT TT-3’; E-box1 mut FW, 5’-AAA GTC ATT Cgc TAg GGC CAG GGG CG-3’, and E-box1 mut RV, 5’-CGC CCC TGG Ccc TAg gga ATG ACT TT-3’. Small letters indicate mutated nucleotides.

**ChIP assay**

Mouse embryos (more than 120 embryos) were dissected at E10.5, and PSMs were collected in ice-cold PBS with protease inhibitor mixture. PSMs were dispersed by 0.05 w/v % trypsin/EDTA treatment for 3 min. Cells were fixed with 1% formaldehyde in PBS for 10 min at room temperature, and the fixation was stopped by adding 0.1 amount of 1.5 M glycerol. After a brief centrifugation, cells were lysed with 200 ml of SDS lysis buffer (50 mM Tris-HCl, pH 8.0, 10 mM EDTA, 1% SDS). Cells lysates were sonicated using a bioruptor (Cosmo Bio) with power set at high and a 30-s on and 60-s off interval. After centrifugation to remove the insoluble fraction, supernatants were incubated with 15 μl of 30% slurry anti-Myc or anti-FLAG (monoclonal, FLA-1, MBL, Japan) overnight, followed by incubation with anti-mouse IgG conjugated with horseradish peroxidase. Signals were visualized by the enhanced chemiluminescence detection system according to the manufacturer's instruction.

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**Promoter elements for restricted expression of Hes7**

HEK293T cells were seeded at 4 × 10⁵ cells in a 10-cm dish, cultured for 24 h, and transfected with 5 μg of each expression vectors or empty pcDNA3 vector. For 48 h of culture, cells were lysed in ice-cold TNE buffer (20 mM Tris-HCl, pH 8.0, 0.1 mM EDTA, 1% SDS). Cell lysates were sonicated using a bioruptor (Cosmo Bio) with power set at high and a 30-s on and 60-s off interval. After centrifugation to remove the insoluble fraction, supernatants were incubated with 15 μl of 30% slurry anti-Myc or anti-FLAG (monoclonal, FLA-1, MBL, Japan) overnight, followed by incubation with anti-mouse IgG conjugated with horseradish peroxidase. Signals were visualized by the enhanced chemiluminescence detection system according to the manufacturer's instruction.

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