Supplemental Information

Acute Induction of *Eya3* by Late-Night Light Stimulation Triggers
*TSHβ* Expression in Photoperiodism

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Supplemental Results and Discussion

Circadian Oscillators in the Mouse PT
To investigate whether PT region contains a circadian oscillator, we examined brain slices of *mPer2* luc mice [1] containing PT and found that the slice exhibited several circadian oscillations of bioluminescence (Figure S1A). Also, it was reported that PER2::LUC rhythms were detected in the median eminence/pars tuberalis [2]. This result and previously report indicate that the PT contains the circadian oscillator.

Circadian Oscillations of Clock and Clock-Controlled Genes in the Mouse PT
Many known clock and clock-controlled genes exhibited 24 hr rhythmicity in the PT under both short-day and long-day conditions (Figure 1B and Table S2). However, some clock and clock-controlled genes are not in the list of 24 hr rhythmic genes. Therefore, we checked 24 hr rhythmicity of known clock and clock-controlled genes in the short-day and long-day conditions by using stringent criteria for 24 hr rhythmicity (correlations > 0.8 both in short-day and long-day conditions). *Per1*, *Clock* and *Tef* exhibited stringent 24 hr rhythmicity in the short-day condition (correlations > 0.8), but not in the long-day condition (correlations < 0.8). *Cry2* and *Hlf* did not exhibit stringent 24 hr rhythmicity either in the short-day or long-day conditions.

Expression of Photoperiodic Gene Candidates in the Mouse PT
The identified photoperiodic genes included *TSHβ*. In contrast to the β subunit of TSH (*TSHβ*), its α subunit (*Cga*) did not respond to the photoperiod and was not identified as a photoperiodic gene by our statistical analysis (Figure S1B). Recently, tachykinin precursor 1 (*Tacl*), which encodes the tachykinin peptide hormone family members substance P and neurokinin A, was
identified as a long-day gene in the sheep PT, and Tac1 has been suggested to regulate prolactin release in vivo [3]. However, the Tac1 gene was not identified as a photoperiodic gene by our statistical analysis (Figure S1B), and this finding was confirmed using quantitative PCR (qPCR) (Figure S1C).

**Expression Dynamics of Eya3 and TSHβ**
The induction of Eya3 was acute and transitory, whereas TSHβ expression was gradually induced. We consider that the difference between time-scale of Eya3 and TSHβ inductions is likely due to a molecular cascade of transcriptional regulation of Eya3 followed by that of TSHβ. In general, a cascade of transcriptional regulation can generate the delay in transcription [4]. Therefore, the observed gradual induction of TSHβ after the acute induction of Eya3 could be, at least in part, explained by the cascade of transcriptional regulation.

**Activation of TSHβ Promoter by Eya3**
In Figure 4B, Eya3, which is a transcriptional co-activator and does not directly bind to DNA, appears to activate the TSHβ promoter when transfected alone. We thus measured Six1 mRNA expression in NIH3T3 cells by qPCR method and found that Six1 is expressed in the NIH3T3 cells (Figure S2C). Therefore, we speculate that transfection of Eya3 alone may be able to slightly increase the activation of TSHβ promoter through the interaction with endogenous Six1.

**Tef and Hlf Enhance the Eya3-Six1-Dependent Activation of TSHβ Promoter**
Since thyrotroph embryonic factor (Tef) can increase the TSHβ promoter activity [5], we examined the contribution of Tef and its family members, hepatic leukemia factor (Hlf) and Dbp to 7.7-kbp and 0.1-kbp TSHβ promoter. These PAR bZIP transcription factors can activate various promoters via the D box consensus sequences [6, 7]. TSHβ promoter contains at least three functional D boxes upstream of the TSS [5]. We first examined the ability of Tef, Hlf, and Dbp to trans-activate the 7.7-kbp or 0.1-kbp TSHβ promoter. We found that the transfection of Tef, Hlf or Dbp significantly increased the Luciferase activity of the 7.7-kbp or 0.1-kbp TSHβ promoter (Figure S2D, P(TSHβ-7.7k) and P(TSHβ-0.1k)), whereas neither activated the SV40 promoter activity, indicating that the Tef-, Hlf- or Dbp-dependent activation of the TSHβ promoter was specific (Figure S2D, P(SV40)). Consistent with these results, the 0.1-kbp TSHβ promoter possesses one of the previously identified D boxes [5], which is highly conserved among vertebrates (Figure 4C).

We note that Dbp slightly decrease Luciferase activity of the 7.7-kbp TSHβ promoter in combination with the Tef or Hlf. This result could be simply explained by the weaker activation of 7.7-kbp TSHβ promoter by Dbp alone than that by Tef or Hlf (Figure S2D, P(TSHβ-7.7k)). A
weaker transcriptional activator (e.g. \textit{Dbp}) apparently serves as a transcriptional repressor by partially inhibiting the binding of a stronger transcriptional activator (e.g. \textit{Tef} or \textit{Hlf}). This hypothesis could be confirmed when we use the 0.1-kbp \textit{TSHβ} promoter, to which \textit{Dbp}, \textit{Tef} or \textit{Hlf} activate almost equally. As predicted, we confirmed that \textit{Dbp} does not decrease \textit{Luciferase} activity of 0.1-kbp \textit{TSHβ} promoter even in combination with the \textit{Tef} or \textit{Hlf} (Figure S2D, \textit{P}(\textit{TSHβ}-0.1k)).

In addition to \textit{Tef}, \textit{Hlf}, and \textit{Dbp}, we also examined \textit{E4bp4} (\textit{Nfil3}), which is also rhythmically expressed in the PT (Figure 1B) because \textit{E4bp4} can function as either a trans-repressor or trans-activator of transcription in different cell types and promoter contexts [8, 9]. We found that \textit{E4bp4} exhibited no significant effect on 0.1-kbp \textit{TSHβ} promoter activity (Figure S2E).

The temporal expression profiles of \textit{Tef}, \textit{Hlf}, and \textit{Dbp} under the long-day and short-day conditions showed that \textit{Tef} and \textit{Dbp} exhibited 24 hr rhythmicity under both the long-day and short-day conditions. In contrast, \textit{Hlf} exhibited only slight 24 hr rhythmicity under the short-day condition and a photoperiodic response under the long-day condition (Figure S2F).

We next investigated the spatial expressions of \textit{Tef}, \textit{Hlf} and \textit{Dbp} genes in the mouse PT under the long-day condition by RI \textit{in situ} hybridization (Figure S2G) and found that the \textit{Tef} and \textit{Hlf} mRNAs were highly expressed whereas the \textit{Dbp} mRNA was expressed only weakly. Based on these findings, we hereafter focused on the contribution of \textit{Tef} and \textit{Hlf} to the \textit{Eya3-Six1}-dependent \textit{TSHβ} promoter activity. When co-transfected with \textit{Eya3} and \textit{Six1}, \textit{Tef} or \textit{Hlf} synergistically increased the \textit{Luciferase} activity of the \textit{TSHβ} promoter (Figure S2H, \textit{P}(\textit{TSHβ}-0.1k)). We also confirmed that \textit{Eya3} increased the \textit{TSHβ} promoter activity in a dose-dependent manner when it was co-expressed with \textit{Six1} and \textit{Tef} or \textit{Hlf} (Figure S2I, \textit{P}(\textit{TSHβ}-0.1k)).

To determine which cis-element(s) in the \textit{TSHβ} promoter (\textit{P}(\textit{TSHβ}-0.1k)) were activated by \textit{Tef} and \textit{Hlf}, we mutated the D box as well as the putative Six consensus sequences in the \textit{TSHβ} promoter, and then performed \textit{Luciferase} transfection assays. We found that the mutation of the D box site in the \textit{TSHβ} promoter significantly decreased the change elicited by the \textit{Tef}- or \textit{Hlf-dependent} activation (Figure S2J, \textit{P}(\textit{TSHβ}-mutD)). In contrast, neither the mutation nor deletion of the MEF3 site had much effect on the \textit{Tef}- or \textit{Hlf-dependent} activation of the \textit{TSHβ} promoter. We also noted that the mutation of the D box site in the \textit{TSHβ} promoter did not affect the change elicited by the \textit{Eya3-Six1-dependent} activation (Figure 4E, \textit{P}(\textit{TSHβ}-mutD)).

These results indicate that \textit{Tef} and \textit{Hlf} act additively on only one D box in the 0.1-kbp \textit{TSHβ} promoter (Figure S2D and J). Since both \textit{Hlf} and \textit{Tef} act as a transcriptional activator of D box, transfection of both \textit{Hlf} and \textit{Tef} into the cells seems equivalent to the increased concentration of a transcriptional activator of D box (either \textit{Hlf} or \textit{Tef}). A transcriptional
activator at higher concentration can bind to a cis-element with higher probability, which will result in the increased transcriptional activity. This simple hypothetical mechanism will explain why co-transfection of both Hlf and Tef additively activates TSHβ promoter containing only one D box.

This D box is located near the transcription start site in the TSHβ promoter, and looks like and might work as a TATA box. To exclude this possibility, we examined the basal activity of the D box-mutated TSHβ promoter (P(TSHβ-mutD)) and compared it with the wild type TSHβ promoter (P(TSHβ-0.1k)). In fact, the basal activity of P(TSHβ-mutD) in Figure 4E was 1.16 ± 0.070-fold (average ± SEM, n = 3) of that of wild type P(TSHβ-0.1k), suggesting that a TATA-like sequence overlapping with D box in the TSHβ promoter does not serve as a TATA box.

We next examined the effect of deleting either the So1 or So2 site in the TSHβ promoter (Figure S2J). Deletion of the So1 site slightly decreased the change elicited by the Tef- or Hlf-dependent activation (Figure S2J). These results suggested that the D box is important for the activation of the TSHβ promoter by Tef or Hlf and that the So1 site also, but slightly, affects the activation of the TSHβ promoter by Tef or Hlf. We also note that both the D box and the So1 site are important for the Eya3-Six1-Tef- or Eya3-Six1-Hlf-dependent activation of the TSHβ promoter (Figure S2J). Deletion of So2 site increased the activation of TSHβ promoter by co-transfection with Eya3, Six1 and Tef. Therefore, So2 site may play a suppressing role in activation of TSHβ promoter although it is unknown what molecular mechanism functions.

Since the So1 and D box are close to one another in the TSHβ promoter, and since specific combinations of neighbouring cis-elements can exhibit qualitatively new properties from those of either element alone [10], we speculate that the Six consensus sequence and D box may constitute a composite promoter and provide combinatorial regulation in the photoperiodic response. Consistent with this hypothesis, transcriptional activators of the Six consensus sequence (Eya3 and Six1) and of the D box (Tef and Hlf) exhibited synergistic activation of the TSHβ promoter. In addition, deletion of the Six consensus sequence affected the Tef- or Hlf-dependent activation of the TSHβ promoter (Figure S2J). In the mouse PT, the Tef and Hlf mRNAs exhibited circadian expression, peaking in the subjective morning under the short-day condition (Figure S2F). Therefore, we expect that the late-night light stimulation induces the expression of Eya3 in the PT with timing that ensures that it is co-expressed with Tef or Hlf during the subjective morning, to synergistically trigger TSHβ expression.

CBA/N Mice as a Model System for the Study of Photoperiodism
This study provided the first genome-wide expression analysis of the CBA/N mouse PT. Our findings in this study, together with the previous report [11] by Dr. Yoshimura in Nagoya
University, suggest that a photoperiodic response of $TSH\beta$ and much of the photoperiodic machinery are preserved in the PT of the CBA/N mouse. These studies have revealed the usefulness of CBA/N mouse as a model organism for the molecular study of photoperiodic system.

**Retinal Degeneration in CBA Mice**

It was reported that CBA/J ($rd/rd$) mice have lower sensitivity to light due to the retinal degeneration, whereas CBA/N (+/+ ) mice have normal light sensitivity [12, 13]. In this study, we used CBA/N mice (+/+ ), which have normal retina.
Supplemental Figure 1. Masumoto et al.
Figure S1. Expression of Some Photoperiodic and Nonphotoperiodic Genes
(Related to Figure 1)
(A) Representative data of bioluminescence showing circadian profiles of mPER2 expression from PT region from mPer2Luc knockin mouse.
(B) Expression of known photoperiodic and nonphotoperiodic genes under the short-day and long-day conditions was measured by GeneChip. The GeneChip data for TSHβ, Cga, and Tac1 mRNA under the long-day and short-day conditions were indicated by a line plot (n = 2). The vertical axis indicates gene expression on a logarithmic scale (base is 2). Error bars represent ± SEM.
(C) Confirmation of the GeneChip data for the Tac1 expression. Tac1 expression under the long-day and short-day conditions measured by qPCR (n = 2). Relative expression to Tbp expression is plotted.
(D) GeneChip data of TSHβ, Eya3, Rorβ, and Maff expression on the first long-day (advance and delay conditions) were indicated by a line plot (n = 2). The vertical axis indicates gene expression on a logarithmic scale (base is 2). Error bars represent ± SEM.
Supplemental Figure 2. Masumoto et al. (1/3)
Supplemental Figure 2. Masumoto et al. (3/3)
Figure S2. The Activities and the Expression Profiles of the Transcriptional Regulators of the TSHβ Promoter (Related to Figure 4)

(A and B) RORβ and MAFF had almost no effect on the TSHβ promoter activity. The TSHβ promoter (-7.7k) was barely activated by RORβ (A) or MAFF (B).

(C) mRNA expression levels of endogenous TSHβ, Eya3 and Six1 in NIH3T3 cells. Relative mRNA levels of each gene to Tbp expression were measured with quantitative PCR assay (n = 3, error bars means ± SEM).

(D) The TSHβ promoter (-7.7k and -0.1k) could be activated by TEF, HLF, and DBP.

(E) The TSHβ promoter (-0.1k) was barely activated by E4BP4 (Nfil3).

(F) Gene expression of Tef, Hlf, and Dbp. GeneChip data for the Tef, Hlf, and Dbp expression under the short-day and long-day conditions and on the first long-day (advance and delay conditions) indicated by line plot (n = 2). The vertical axis indicates gene expression on a logarithmic scale (base is 2). Error bars means ± SEM.

(G) Tef, Hlf, and Dbp expression in the mouse PT at ZT8 under the long-day condition, detected by R1 in situ hybridization. Scale bar=300 μm.

(H) Tef and Hlf enhance the Eya3-Six1-dependent activation of TSHβ promoter. The TSHβ promoter (-0.1k) was strongly activated by EYA3, SIX1, and TEF or HLF. Experiments of Figure 4B and Figure S2H were performed at the same time, so Luciferase activities of P(TSHβ-0.1k) and P(SV40) in Figure 4B were re-plotted in Figure S2H as control data.

(I) TSHβ expression is synergistically activated by Eya3 and Six1, and further enhanced by Tef and Hlf.

(J) Role of the D box and So1 site in transcriptional activation of the TSHβ promoter by Eya3, Six1, and Tef or Hlf. (Top panel) The TSHβ promoter (-0.1k), and its MEF3-deleted (P(TSHβ-ΔMEF3)) and MEF3-mutated (P(TSHβ-mutMEF3)) forms were strongly activated by EYA3, SIX1, and TEF or HLF. The D box-mutated TSHβ promoter (P(TSHβ-mutD)) was not activated by TEF and HLF. (Bottom panel) The So2-deleted form (P(TSHβ-ΔSo2)) were strongly activated by EYA3, SIX1, and TEF or HLF. The So1-deleted TSHβ promoter (P(TSHβ-ΔSo1)) was not strongly activated.

(A, B, D, E, H-J) Each of the indicated promoters was fused to a Luciferase reporter gene and used to transiently transf ect NIH3T3 cells. The Luciferase activity for each promoter is expressed relative to the activity with an empty vector. Data are representative of two independent experiments. Error bars means ± SEM. (n = 3).
Supplemental Tables (see accompanying Excel spreadsheet)

Table S1. List of 24 hr Rhythmic Genes in the Mouse PT under Short-Day and Long-Day Conditions (Related to Figure 1A)
The table provides the Affymetrix probe set ID, p value, FDR, mean of the phases under short-day and long-day conditions, phase difference between short-day and long-day conditions, phase in the short-day condition, phase in the long-day condition, the best Pearson's correlation to cosine curves of a 24 hr period in short-day condition, the best Pearson's correlation to cosine curves of a 24 hr period in long-day condition, amplitude of circadian expression in short-day condition, amplitude in long-day condition, relative amplitude in short-day condition, p value in short-day condition, p value in long-day condition, description of the probe set, gene name, and gene symbol.

Table S2. List of Clock and Clock-Controlled Genes with Significant Circadian Expression under Short-Day and Long-Day Conditions (Related to Figure 1B)
The table provides the gene symbol, Affymetrix probe set ID, phase in short-day condition, the best Pearson's correlation to cosine curves of a 24 hr period in short-day condition, phase in long-day condition, the best Pearson's correlation to cosine curves of a 24 hr period in long-day condition, the minimum of the values in the 4th and 6th columns (the smaller value of two best Pearson's correlation), and difference between the phases in short-day and long-day conditions.

Table S3. List of Long-Day and Short-Day Genes under Chronic Short-Day and Long-Day Conditions (Related to Figure 1C)
The table provides the Affymetrix probe set ID, p value, FDR, q value, whether the expression in long-day condition is up-regulated ('up') or down-regulated ('down') against the short-day condition, fold change between the mean expression in short-day and long-day conditions, description of the probe set, gene name, and gene symbol.

Table S4. List of Acute Long-Day Genes (Related to Figure 3A)
The table provides the Affymetrix probe set ID, p value, FDR, q value of difference between short-day and advance conditions, maximum fold change between the lowest and highest mean expressions in the short-day and advance conditions, p value, FDR, q value of difference between short-day and delay conditions, maximum fold change between the lowest and highest mean expressions in the short-day and delay conditions, description of the probe set, gene name, and gene symbol.
Supplemental Experimental Procedures

Circadian-Time Analysis from Time-Course Expression Data
To detect how much the circadian phase differed between the short-day and long-day conditions, we investigated the expressions of the following 20 clock and clock-controlled genes by GeneChip: Bmal1 (Arntl), Bmal2 (Arntl2), Clock, Cry1, Cry2, Dbp, Dec1 (Bhlhb2), Dec2 (Bhlhb3), E4bp4 (Nfil3), Npas2, Per1, Per2, Per3, Tef, Hlf, Rev-Erbα (Nr1d1), Rev-Erbβ (Nr1d2), Rora, Rorβ, and Rory. We used natural expression values from the GeneChip data. We concatenated two time-courses of experimental replicates (6 time points for each replicate) and used 12 time-point dataset for the subsequent analysis. For each gene, we calculated the best Pearson's correlation between gene expression data and cosine curves of a 24 hr period, and determined its phase in short-day and long-day conditions using a Fourier transformation-based method [14]. If the GeneChip had multiple probe sets for a single gene, we first calculated, for each probe set, two best Pearson's correlation from short-day and long-day conditions, respectively, and then registered the smaller best Pearson's correlation. We selected the probe set with the maximum value of the registered (i.e. smaller) best Pearson's correlation. Next, we chose genes for which the best Pearson's correlation for both the short-day and long-day conditions was more than or equal to 0.8 (see Figure 1B). Finally, we calculated the median of the phase difference (long-day – short-day) for these RNAs. We used this median value (~7.71 hours) as the phase delay of the long-day versus the short-day condition.

Identification of 24 hr Rhythmic Genes
As described above, we concatenated two time-courses of experimental replicates (6 time points for each replicate) and used 12 time-point dataset for the subsequent analysis. To identify 24 hr rhythmic genes in the mouse PT, we first calculated the best Pearson's correlation between gene expression data and cosine curves of a 24 hr period, and determined its phase in short-day and long-day conditions using a Fourier transformation-based method [14]. We then calculated the p values for the best correlations of the short-day and long-day using random shuffling, and then combined these two p values with Fisher's method \( P = P_1P_2\left[1 - \log(P_1P_2)\right] \). We estimated the false-discovery rates (FDRs) of the combined p value based on the Benjamini-Hochberg method [15], and selected significantly oscillating genes as those for which the FDRs were less than or equal to 0.05.

Identification of Photoperiodic Genes
To identify the photoperiodic genes, we used time-course expression data under chronic short-day and long-day conditions. In order to complement the observed phase difference
between short-day and long-day conditions (the approximately 8 hours, see above), we shifted time points by 8 hours; that is, the short-day ZT16, 20, 0, 4, 8, 12 corresponded to long-day ZT0, 4, 8, 12, 16, 20, respectively. We used log-2-transformed expression values of the GeneChip data in the following analysis. We performed a two-way ANOVA to test the difference in expression levels between the short-day and long-day conditions for each probe set, and assigned p values and FDRs. We selected as significantly differently expressed genes those whose FDRs were less than or equal to 0.05 and for which the difference between the maximum expression value in the short-day condition and the minimum value in the long-day condition (“short-day” gene) or between the maximum expression in the long-day condition and the minimum in the short-day condition (“long-day” gene) was more than 2.5-fold. In this method, we did not subtract the 24 hr rhythmic genes that were identified above, and thus 38 genes were identified both in the 24 hr rhythmic and the photoperiodic genes.

Circadian-Time Estimation from Single-Time-Point Expression Data
To estimate how the circadian phases at ZT16 were changed on days 0, 1, 3 and 5 after transition from the short-day to the delay condition, we used a previously reported molecular-timetable method [16, 17]. To construct a molecular timetable that has the phase (peak time), mean, and standard deviation (SD) of the gene expressions of clock and clock-controlled genes, we used the expression values of 20 clock and clock-controlled genes from the qPCR data at every 4 hours during 48 hours in the short-day condition. We first normalized the qPCR expression data of the 20 clock and clock-controlled genes by using Tbp expression level at the same time point. We then calculated the best Pearson's correlation between cosine curves of a 24 hr period and the qPCR expression data. We selected genes with the best Pearson's correlation ≥ 0.8. For these genes, we also calculated peak time, mean and SD of the qPCR expression data. With this molecular timetable, we determined the circadian phase of the mouse PTs at ZT16 on days 0, 1, 3, and 5. In details, we normalized each qPCR expression value obtained at ZT16 for each day by subtracting the estimated mean and then divided by the SD in the molecular timetable. We then plotted the data on a graph, with the phase (peak time) on the x-axis and normalized expression values on the y-axis. From this graph, we searched for the cosine curve, $y = \sqrt{2} \times \cos(x-b)$, that had the maximum correlation, and determined $b$ as the phase of the day.

Identification of Acute Long-Day Genes
To identify acute long-day genes whose expression patterns were changed on the first long-day after the transition from the short-day to the long-day condition, we sought to retrieve probe sets whose expression patterns in the short-day condition and advance condition were significantly different but whose expression patterns in the short-day condition and delay condition were not
significantly different. First, we estimated that the phase delay of the advance condition and the delay condition against the short-day condition was approximately 0 hours (~0.88 hours), and 4 hours (~3.96 hours), respectively, using the method described in the above circadian-time analysis from the time-course expression data. In order to complement these phase differences, we set the short-day ZT0 to correspond with the advance condition ZT0, and the short-day ZT20 with the delay condition ZT0. We then performed the same statistical procedures described above for identifying genes whose time-course expression patterns in the short-day and long-day condition were different. In this case, we chose genes whose expressions were significantly different between the short-day condition and the advance condition, but were not significant between the short-day condition and the delay condition.

**Preparation of PT Region Slices from mPer2Luc Mice**

The brains were removed from quickly decapitated, young (older than 4 weeks of age) mPer2Luc mice, then 300-μm thick coronal sections containing the PT were made using a vibratome type linear slicer (PRO7; Dosaka EM Corp) in ice cold Hanks' balanced salt solution (Invitrogen). PT region slices included a small amount of the surrounding tissue. The slices were then placed on a culture membrane (MilliCell-CM; Millipore) and set on a dish with 1.2 mL culture medium [DMEM supplemented with 1.2 g/L NaHCO3 (Nacalai Tesque), 15 mM HEPES (Dojindo), 20 mg/L kanamycin (Invitrogen), 5 μg/mL insulin (Sigma), 20 nM putrescine (Sigma), 100 μg/mL apo-transferrin (Sigma), 20 nM progesterone (Sigma), 30 nM sodium selenite (Sigma), one-fiftieth part B-27 supplement (Invitrogen), and 1 mM luciferin]. The culture dishes were placed in a high-sensitivity bioluminescence detection system (LM-2400; Hamamatsu Photonics), and the bioluminescence of each well was measured at 37°C.

**Quantitative PCR (qPCR) of PT Samples**

The total RNA was prepared from the pooled PT samples obtained at each time point under each condition, using Trizol reagent (Gibco BRL). qPCR was performed with the ABI Prism 7900 and SYBR Green Reagents (Applied Biosystems). The cDNAs were synthesized from 0.25 μg of total RNA using Superscript II reverse transcriptase (Invitrogen). Samples contained 1×SYBR Green Master Mix, 0.8 μM primers, and 1/40 synthesized cDNA in a 10 μl volume. The PCR conditions were as follows: 10 min at 95°C, then 45 cycles of 15 s at 94°C, 1 min at 59°C. The absolute cDNA abundance was calculated using a standard curve obtained from murine genomic DNAs. We used Tbp as the internal control.

**Oligonucleotide Sequences for qPCR of Tissue Samples**

The primers (Hokkaido System Science) used in qPCR were as follows:
In Situ Hybridization (ISH)
Mice were deeply anesthetized with ether and intracardially perfused with 10 ml saline and 20 ml of a fixative containing 4% paraformaldehyde in 0.1 M phosphate buffer (PB), pH 7.4. Mouse brain samples were postfixed in the same fixative for 24 hours at 4°C, soaked in PB containing 20% sucrose for 48 hours, and stored frozen at -70°C. The ISH method was described in detail previously [18]. Serial coronal sections (40-µm thick) of the mouse brain were made using a cryostat. Fragments of cDNA were obtained by PCR, and the products were subcloned into the pGEM-T easy vector (Promega). Radiolabelled probes were generated using 35S-UTP (PerkinElmer) via a standard protocol for cRNA synthesis. The probe sequence information is presented in the Supplemental Experimental Procedures.

Oligonucleotide Sequences for ISH
The primers (Hokkaido System Science) used in the construction of ISH cRNA probes were as follows:

**TSHβ cRNA probe:**
- **Forward primer:** 5'-TGGGTGGAGAAGAGTGAGCG-3'
- **Reverse primer:** 5'-ACCAGATTGCACTGCTATTG-3'

**Eya1 cRNA probe:**
- **Forward primer:** 5'-GGACTTTTACTACATAACTGGGTCAGCAGC-3'
- **Reverse primer:** 5'-GGTCTCAACACTGGCCATTTCTGTCTG-3'

**Eya2 cRNA probe:**
- **Forward primer:** 5'-GAATCATGCAGAGGTTTGGCCGCAAAG-3'
- **Reverse primer:** 5'-CTCCCTTGCCAATGCCCAGAGAATGATTG-3'
**Eya3 cRNA probe:**
- Forward primer: 5'-AGCACAATGCCAGCCTGATAACCAC-3'
- Reverse primer: 5'-CTGTTGGGTCCTTTCCATACTTCTG-3'

**Eya4 cRNA probe:**
- Forward primer: 5'-TGCCAACAGGTGTGAGGAGGAGG-3'
- Reverse primer: 5'-CCAGTGCTTGAGGAGGAATTCTC-3'

**Six1 cRNA probe:**
- Forward primer: 5'-CCTTCCTCCTCGTCTTTTAATGAACAGA-3'
- Reverse primer: 5'-CGATGGCAGCCTGCTGAGGAGTTTCTG-3'

**Six2 cRNA probe:**
- Forward primer: 5'-ACATCGAGCGGCTGGGCCGCTTCCTG-3'
- Reverse primer: 5'-CGATGGCAGCCTGCTGAGGAGTTTCTG-3'

**Six3 cRNA probe:**
- Forward primer: 5'-CCTCCTCCTCGTCTTTTAATGAACAGA-3'
- Reverse primer: 5'-CGATGGCAGCCTGCTGAGGAGTTTCTG-3'

**Six4 cRNA probe:**
- Forward primer: 5'-TGGTCATGACATGGGGCATGAGTCAAAAC-3'
- Reverse primer: 5'-GGGGTCAGGGAGTCATTTAAGACCCGGAG-3'

**Six5 cRNA probe:**
- Forward primer: 5'-TGACAATGGTGAGGTGACAGGAGG-3'
- Reverse primer: 5'-GGGGTCAGGGAGTCATTTAAGACCCGGAG-3'

**Six6 cRNA probe:**
- Forward primer: 5'-CATCCAGCGACAGTGAGTGAG-3'
- Reverse primer: 5'-CATAGCAGCACTGTCATGCGGCGTTCCAC-3'

**Dach1 cRNA probe:**
- Forward primer: 5'-TGGTGTCATGACATGGGGCATGAGTCAAAAC-3'
- Reverse primer: 5'-GGGGTCAGGGAGTCATTTAAGACCCGGAG-3'

**Dach2 cRNA probe:**
- Forward primer: 5'-GAGGGGTCAGGGAGGTGACAGGAGG-3'
- Reverse primer: 5'-GAGGGGTCAGGGAGGTGACAGGAGG-3'

**Ski cRNA probe:**
- Forward primer: 5'-AGATACCTCAGGAGAGGTGAGGAGGG-3'
- Reverse primer: 5'-AGATACCTCAGGAGAGGTGAGGAGGG-3'

**Skil cRNA probe:**
- Forward primer: 5'-AGATACCTCAGGAGAGGTGAGGAGGG-3'
- Reverse primer: 5'-AGATACCTCAGGAGAGGTGAGGAGGG-3'
Dbp cRNA probe:
- **Forward primer:** 5'-ACCGCGCAGGCTTGACATCTAGGGAC-3'
- **Reverse primer:** 5'-CATGACGTTCCTCGGCACCTAGCTGG-3'

Tef cRNA probe:
- **Forward primer:** 5'-TGGAAGAGCCACCCTTTGGAGGAC-3'
- **Reverse primer:** 5'-CACACACATGTAAGGCCAAAGGGCTGC-3'

Hlf cRNA probe:
- **Forward primer:** 5'-GTGCATGACCAGAATATTCTCAGGACAGGG-3'
- **Reverse primer:** 5'-CGGAGAAACCACAGTTTGAGACAACACC-3'

qPCR of NIH3T3 Cell Samples
NIH3T3 cells were grown in DMEM (Invitrogen) supplemented with 10% FBS (JRH Biosciences) and antibiotics (100 U/ml penicillin and 100 µg/ml streptomycin; Invitrogen). Total RNA was prepared from NIH3T3 cells using TRiZol reagents (Invitrogen) according to the manufacturer's instructions. The cDNA was synthesized from 0.25 µg of total RNA with random 6 mer (Promega) and Superscript II Reverse Transcriptase (Invitrogen) according to the manufacturer's instructions. qPCR was performed using ABI Prism 7900 and Power SYBR Green Reagents (Applied Biosystems). Samples contained 1×Power SYBR Green Master Mix (Applied Biosystems), 0.8 µM primers and 1/40 synthesized cDNA in a 10 µl volume. PCR conditions were as follows: 10 min at 95°C, then 45 cycles of 15 s at 94°C and 1 min at 59°C. Absolute cDNA abundance was calculated using a standard curve obtained from murine genomic DNA. *Tbp* expression levels were quantified and used as an internal control.

Oligonucleotide Sequences for qPCR
The primers (Hokkaido System Science) used in qPCR were as follows:

**TSHβ mRNA:**
- **Forward primer:** 5'-CTGCATACACGAGGCTGCAG-3'
- **Reverse primer:** 5'-CCCCAGATAGAAAGACTGCGG-3'

**Eya3 mRNA:**
- **Forward primer:** 5'-TTCACAGCTCCAAGTAGAATCTGACT-3'
- **Reverse primer:** 5'-TATGGAAGCGCCATGAGCTT-3'

**Six1 mRNA:**
- **Forward primer:** 5'-TATTTTTGAAGTCTGTCACCCGAA-3'
- **Reverse primer:** 5'-TTCGTCATATCATTAACCTAGCCACT-3'
**Tbp mRNA:**

**Forward primer:** 5'-CCCCCTCTGCACTGAAATCA-3'

**Reverse primer:** 5'-GTAGCAGCAGAGCAAGCAA-3'

**Transfection and Luciferase Assay**

NIH3T3 cells (American Type Culture Collection) were maintained in DMEM (Invitrogen) supplemented with 10% FBS (JRH Biosciences) and antibiotics (100 U/ml penicillin and 100 µg/ml streptomycin; Invitrogen). One day prior to transfection, the cells were plated onto 35-mm dishes at a density of 2×10^5 cells per dish. The following day, the cells were co-transfected using FuGene6 (Roche) with 0.4 µg of a Luciferase reporter plasmid in the presence of the following constructs, as indicated in the respective figures: 0 or 0.4 µg of pCMV-TnT-Rorβ (for Figure S2A); 0 or 0.4 µg of pCMV-SPORT6-Maff (MGC clone # 30560, Invitrogen, for Figure S2B); 0 or 0.4 µg each of pMU2-Eya3 and pMU2-Six1 (for Figure 4B, E and F); 0, 0.133, 0.4 or 1.2 µg of pMU2-Eya3 and 0 or 0.4 µg of pMU2-Six1 (for Figure 4D); 0 or 0.4 µg each of pMU2-Dhp [19], pMU2-Tef and pMU2-Hlf (for Figure S2D); 0 or 0.4 µg each of pMU2-Tef, pMU2-Hlf and pMU2-E4bp4 [19] (for Figure S2E); 0 or 0.4 µg each of pMU2-Eya3, pMU2-Six1, pMU2-Tef and pMU2-Hlf (for Figure S2H and J); 0, 0.133, 0.4 or 1.2 µg of pMU2-Eya3 and 0 or 0.4 µg of pMU2-Six1, pMU2-Tef and pMU2-Hlf (for Figure S2I). Empty vector was used to bring the total amount of DNA to 4.0 µg per well. In addition, 50 ng of a phRL-SV40 plasmid (Renilla Luciferase (RLuc) reporter vector, Promega) was added to each transfection as an internal control for transfection efficiency. pGL4.13 (Promega) was used for the construction of the control promoter (P(SV40)). Forty-eight hours after the transfection, the cells were harvested and assayed with the Dual-Luciferase Reporter Assay System (Promega). The Luciferase activity was normalized to the RLuc activity.

**Construction of Plasmids**

**pGL4.10-P(TSHβ-7.7k)**

A 7.7-kbp promoter region of the mouse TSHβ gene was amplified from C57BL/6 genomic DNA using PCR with the following primers (Hokkaido System Science, restriction recognition sequences are underlined): Forward primer containing the XhoI recognition sequence, (5'-AAGCCTCGAGGTTGGTCTGGGATGACTGCTAAGA-3'); reverse primer containing EcoRV recognition sequence, (5'-TTCTGATATCGTGGACACCTACCTTTACCTGCTGAGTG-3'). The PCR product was digested with XhoI and EcoRV, and cloned into the XhoI-EcoRV site of a pGL4.10 vector (Promega) immediately upstream of Luc2, to obtain pGL4.10-P(TSHβ-7.7k). TSHβ promoter constructs were cloned at the +30 position because there is an MEF3 site at +1 position of TSHβ
gene region (Figure 4C). We thus designed TSHβ promoter constructs so that it covers a sufficient flanking sequence of the MEF3 site.

pCMVTnT-Rorβ
The full-length coding sequence of mouse Rorβ was amplified from an NIH3T3 cDNA library using PCR with the following primers: Forward primer, 5'-ATGCGAGCACAAATTGAAGTGATACC-3' (Hokkaido System Science); reverse primer, 5'-TCATTTCAGACCGCAGCACAGTCAGG-3'. The PCR product was treated with the Mighty Cloning kit (TaKaRa) for 5'-end phosphorylation, and cloned into the SmaI sites of a pCMVTnT vector (Promega). The resulting construct was designated as pCMVTnT-Rorβ. In this pCMVTnT vector, the gene is regulated by a CMV promoter.

pGL4.10-P(TSHβ-0.6k), pGL4.10-P(TSHβ-0.2k), and pGL4.10-P(TSHβ-0.1k)
We used pGL4.10-P(TSHβ-7.7k) plasmid DNA as a template for cloning the 0.6- , 0.2- , and 0.1-kbp promoter regions of TSHβ. The 0.6- , 0.2- , and 0.1-kbp TSHβ promoter regions were amplified, treated with the Mighty Cloning kit (TaKaRa) for 5'-end phosphorylation, and cloned into the EcoRV sites of a pGL4.10 vector (Promega). The resulting constructs were designated as pGL4.10-P(TSHβ-0.6k), pGL4.10-P(TSHβ-0.2k), and pGL4.10-P(TSHβ-0.1k).

Oligonucleotide Sequences for Constructing the Shorter TSHβ Promoters (Hokkaido System Science)
Cloning of P(TSHβ-0.6k):
Forward primer: 5'-CCAAGACAGAATTTGATCTCAGGTCAGT-3'
Reverse primer: 5'-CACTCTCCGTTTCATTATTACCTTCA-3'
Cloning of P(TSHβ-0.2k):
Forward primer: 5'-GATATGGTTTTCAATAGAAGAGGAAG-3'
Reverse primer: 5'-CACTCTCCGTTTCATTATTACCTTCA-3'
Cloning of P(TSHβ-0.1k):
Forward primer: 5'-AGATGCTTTTCAGATAAGAAAGCAGC-3'
Reverse primer: 5'-CACTCTCCGTTTCATTATTACCTTCA-3'

pMU2-Eya3 and pMU2-SixI
The full-length coding sequences of mouse Eya3 and Six1 were amplified from the NIH3T3 cDNA library by PCR, using forward primers containing the I-SceI recognition sequence and reverse primers containing the PI-PspI recognition sequence (Hokkaido System Science, restriction recognition sequences are underlined). The PCR products were digested with I-SceI
and PI-PspI (New England BioLabs) and cloned into the pMU2 vector [20], and the resulting vectors were designated as pMU2-Eya3 and pMU2-Six1. In these pMU2 vectors, the genes are regulated by a CMV promoter.

Oligonucleotide Sequences for the Cloning of Eya3 and Six1

Cloning of Eya3:
- **Forward primer:** 5’-ATTACCCCTGTTATCCCTAATGAAGAAGAGCAAGAC CTACCAGAGC-3'
- **Reverse primer:** 5’-ACCCATAATACCCATAATAGCTGTTTGCCATCAGAG GAAGTCAAGCTCTAAAGCC-3'

Cloning of Six1:
- **Forward primer:** 5’-ATTACCCCTGTTATCCCTAATTCGATGCTGCCGTCGT TTGGTT-3'
- **Reverse primer:** 5’-ACCCATAATACCCATAATAGCTGTTTGCCAGGAACC CAAGTCCACCAAACCTG-3'


We used pGL4.10-P(TSHβ-0.1k) as a template. The inverse PCR products were treated with the Mighty Cloning kit (TaKaRa) for 5’-end phosphorylation, and subsequently self-ligated. The constructs were designated as pGL4.10-P(TSHβ-ΔMEF3), pGL4.10-P(TSHβ-mutMEF3), pGL4.10-P(TSHβ-mutD), pGL4.10-P(TSHβ-ΔSo1), and pGL4.10-P(TSHβ-ΔSo2).

Oligonucleotide Sequences for Constructing Mutated or Deleted TSHβ Promoters (Hokkaido System Science)

Inverse PCR of pGL4.10-P(TSHβ-AMEF3):
- **Forward primer:** 5’-AGGGTATAAAATGAACGGAGAGTGG-3'
- **Reverse primer:** 5’-AATCCCCCTCTGATCTTCTTG-3'

Inverse PCR of pGL4.10-P(TSHβ-mutMEF3):
- **Forward primer:** 5’-CATCTAGAGGGTATAAAATGAACGGAGAGTGG-3'
- **Reverse primer:** 5’-AATCCCCCTCTGATCTTCTTG-3'

Inverse PCR of pGL4.10-P(TSHβ-mutD):
- **Forward primer:** 5’-CACCAGCACAAGAAGATCAGAGGGGAATTATCCTG-3'
- **Reverse primer:** 5’-TTGCAATTGAATTGCTGCTTTTATTCTG-3'
Inverse PCR of pGL4.10-P(TSHβ-ASo1):  
**Forward primer:** 5'-GCAATTATATAACAAAGAAGATCAGAGG-3'  
**Reverse primer:** 5'-TGCTTTCTTTATCTGAAAAGCATCTATCC-3'

Inverse PCR of pGL4.10-P(TSHβ-ASo2):  
**Forward primer:** 5'-TTCGAATGCAATTATATAACAAAGAAGATC-3'  
**Reverse primer:** 5'-TTATCTGAAAAGCATCTATCCTCGAG-3'

**Author Contributions**  
K.D.U. performed the GeneChip and qPCR experiments. K.M., M.N., and K.T. performed the ISH experiments. T.K. designed and performed the computational analyses, and constructed the integrated database. M.U. performed the promoter assay. K.M., M.U., T.K., Y.S., and H.R.U. wrote the manuscript. All authors discussed the results and commented on the manuscript.
Supplemental References


