Tissue-specific BMAL1 cistromes reveal that rhythmic transcription is associated with rhythmic enhancer–enhancer interactions

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The mammalian circadian clock relies on the transcription factor CLOCK:BMAL1 to coordinate the rhythmic expression of thousands of genes. Consistent with the various biological functions under clock control, rhythmic gene expression is tissue-specific despite an identical clockwork mechanism in every cell. Here we show that BMAL1 DNA binding is largely tissue-specific, likely because of differences in chromatin accessibility between tissues and cobinding of tissue-specific transcription factors. Our results also indicate that BMAL1 ability to drive tissue-specific rhythmic transcription is associated with not only the activity of BMAL1-bound enhancers but also the activity of neighboring enhancers. Characterization of physical interactions between BMAL1 enhancers and other cis-regulatory regions by RNA polymerase II chromatin interaction analysis by paired-end tag (ChIA-PET) reveals that rhythmic BMAL1 target gene expression correlates with rhythmic chromatin interactions. These data thus support that much of BMAL1 target gene transcription depends on BMAL1 capacity to rhythmically regulate a network of enhancers.

[Keywords: circadian clock; enhancer–enhancer interactions; tissue-specific cistromes; transcription]

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Circadian clocks are found ubiquitously across all kingdoms of life and provide a time-keeping mechanism for organisms to anticipate rhythmic environmental changes. In mammals, virtually every cell harbors the same circadian clockwork that regulates rhythmic gene expression along with temporal cues and systemic signals, such that biological functions occur at the most appropriate time of day. The mammalian circadian clock relies on transcriptional feedback loops initiated by the heterodimeric transcription factor CLOCK:BMAL1 (Partch et al. 2014; Takahashi 2017). CLOCK:BMAL1 rhythmically binds DNA to drive the rhythmic transcription of the genes Period (Per1, Per2, and Per3) and Cryptochrome (Cry1 and Cry2), which, upon translation, form a repressive complex that feedbacks to inhibit CLOCK:BMAL1-mediated transcription. CLOCK:BMAL1 binds to E-boxes and activates transcription more potently during the middle of the day, and maximal repression occurs during the middle of the night (Rey et al. 2011; Koike et al. 2012). CLOCK: BMAL1 is also responsible for the transcriptional regulation of many clock-controlled genes, which allows for rhythms in biochemistry, physiology, and behavior (Panda 2016).

Characterization of the rhythmic transcriptional outputs driven by the circadian clock indicates that genes expressed in a rhythmic manner vary greatly between tissues in the mouse (Panda et al. 2002; Storch et al. 2002; Zhang et al. 2014). Similar findings in plants, insects, and primates also revealed that circadian gene expression is largely tissue-specific (Endo et al. 2014; Meireles-Filho et al. 2017).
However, it is still unknown how the same circadian clock mechanism can generate tissue-specific rhythmic gene expression. Previous work in *Drosophila* revealed that CLOCK:CYC, the CLOCK:BMAL1 homolog, exhibits tissue-specific DNA binding between the body and head, and that much of this tissue specificity is due to the cooperation between CLOCK:CYC and tissue-specific transcription factors [ts-TFs] [Meireles-Filho et al. 2014]. Tissue-specific binding of another mammalian clock component, *Rev-erba*, has also been described, but the underlying mechanisms have not yet been characterized [Zhang et al. 2015].

To address the mechanisms by which the mammalian circadian clock generates tissue-specific circadian transcriptional programs, we performed BMAL1 chromatin immunoprecipitation (ChIP) at the genome-wide level (ChIP-seq) in the mouse liver, kidney, and heart. Our data revealed that the majority of BMAL1 DNA binding is tissue-specific and suggest that accessibility of the chromatin as defined by DNase I-hypersensitive sites [DHSs] [Thurman et al. 2012] and that cobinding of ts-TFs accounts for much of this tissue specificity. In addition, we found large discrepancies between BMAL1 DNA binding and tissue-specific transcription factors with, for example, many genes targeted by BMAL1 in all three tissues only exhibiting rhythmicity in a single tissue. Characterization of the underlying mechanism suggests that the ability of BMAL1 to drive tissue-specific rhythmic gene expression depends on how BMAL1 promotes not only the activity of the DHSs within it but also the activity of other neighboring DHSs. Together, our data suggest that BMAL1 transcriptional output is controlled through enhancer–enhancer interactions and that BMAL1-driven rhythmic transcription depends on the capacity of BMAL1 to rhythmically regulate a network of DHSs.

Results

**BMAL1 DNA-binding signal is largely tissue-specific**

To determine whether BMAL1 cistrome differs between tissues, we performed BMAL1 ChIP-seq in the mouse liver, kidney, and heart with three biological replicates per tissue. Because CLOCK:BMAL1 DNA-binding sites in the liver are virtually identical over the course of the day and only exhibit homogenenous differences in DNA-binding strength [Rey et al. 2011; Koike et al. 2012], we performed BMAL1 ChIP at the time of maximal binding; i.e., Zeitgeber time 6 [ZT6; ZT0 is defined at the light on signal] [Supplemental Fig. S1A]. Identification of BMAL1 DNA-binding sites using HOMER software [Heinz et al. 2010] revealed that most BMAL1 peaks were tissue-specific, with <6% of BMAL1 peaks common to the liver, kidney, and heart [Fig. 1A,B; Supplemental Table S1]. Closer inspection revealed that tissue-specific BMAL1 peaks were more often associated with BMAL1 signal below peak detection threshold in the other tissues rather than a complete absence of signal [Fig. 1B; Supplemental Fig. S1B; see below]. In addition, quantification of BMAL1 ChIP-seq signal revealed that stronger BMAL1 binding is associated with peaks common to more than one tissue [Fig. 1C; see below].

To ensure that differences in BMAL1 ChIP-seq signal between tissues represent true biological variation, we compared BMAL1 ChIP-seq signal between each of the nine data sets and found a high degree of correlation between samples originating from the same tissue but not between tissues [Fig. 1D]. Interreplicates variation between replicates was higher for the kidney than for the liver and heart; however, this might reflect the higher cell type heterogeneity and the variability in the chromatin accessibility landscape that are observed in the kidney when compared with the heart and liver [Cusanovich et al. 2018]. We also found that BMAL1 ChIP-seq signal at core clock genes is nearly identical between all three tissues, further indicating that differences in BMAL1 binding signal between tissues stem from biological variations [Fig. 1E]. Consistent with this finding, gene ontology analysis of BMAL1 target genes for each tissue revealed that while circadian rhythm and other general terms are enriched in all three tissues, tissue-specific biological processes are enriched in a tissue-specific manner [Fig. 1F].

Many of these tissue-specific processes have been shown to be regulated by the circadian clock, such as glycogen synthesis, lipid metabolism, and cholesterol homeostasis in the liver [Doi et al. 2010; Gnocchi et al. 2015; Ma et al. 2015] and blood pressure and vascular smooth muscle contraction in the heart [Wang et al. 2008; Xie et al. 2015]. Together, these results indicate that BMAL1 predominantly binds DNA in a tissue-specific manner and targets genes involved in the regulation of tissue-specific biological functions.

**Chromatin accessibility contributes to tissue-specific BMAL1 DNA binding**

To understand the large differences in BMAL1 ChIP-seq signal between tissues, we set out to define the mechanisms that enable BMAL1 to bind DNA in a tissue-specific manner. Characterization of the chromatin landscape by the ENCODE consortium and others revealed that 94.4% of TF ChIP-seq peaks are located in an accessible chromatin region, i.e., a region that is hypersensitive to DNase I [Thurman et al. 2012; Yue et al. 2014]. Because mouse liver CLOCK:BMAL1 DNA-binding sites are located predominantly in DHSs [Sobel et al. 2017; Trott and Menet 2018], we hypothesized that tissue-specific BMAL1 binding may be, at least in part, caused by differences in DNA accessibility between the liver, kidney, and heart. To test our hypothesis, we used DNase sequencing [DNase-seq] data sets from the mouse ENCODE project [Vierstra et al. 2014; Yue et al. 2014] and found that many tissue-specific BMAL1 peaks are located in a chromatin region that is accessible only in the corresponding tissue [Fig. 2A]. Quantification of DNase-seq signal at liver-, kidney-, and heart-specific BMAL1 peaks [named groups 5, 6, and 7 here, respectively] confirmed that tissue-specific BMAL1 peaks are associated with higher DNase-seq signal [Fig. 2B]. However, some DNase-seq signal could be detected above background level in all three
tissues, suggesting that tissue-specific BMAL1 peaks may not be always associated with a complete absence of DNase-seq signal in other tissues.

To investigate whether tissue-specific BMAL1 binding occurs at DHSs common to several tissues, we mapped the genomic location of DHSs in the liver, kidney, and heart [Supplemental Fig. S2A], and determined the fraction of tissue-specific BMAL1 peaks found at tissue-specific DHSs or at DHSs common to more than one tissue, focusing primarily on liver-specific BMAL1 peaks. As expected, almost all mouse liver BMAL1 peaks (97.7%) are located in chromatin regions that are accessible in the mouse liver (Fig. 2C). However, only ∼20% of liver-specific BMAL1 peaks are located at a liver-specific DHS; i.e., most liver-specific BMAL1 peaks are located within chromatin regions that are also accessible in other tissues (Fig. 2C). Importantly, we found that while no BMAL1 signal is detectable in the kidney and heart at liver-specific DHSs, some BMAL1 signal is visible at common DHSs even if it is not defined as significant BMAL1 peaks by HOMER [Fig. 2C–E]. Similar results were observed for kidney- and heart-specific BMAL1 peaks [Supplemental Fig. S2B, C]. Because DNase I hypersensitivity does not assess the activity of accessible chromatin regions, we also analyzed public ChIP-seq data sets for histone modifications found at transcriptionally active/primed DHSs, i.e., acetylated Lys27 of histone 3 (H3K27ac) and monomethylated Lys4 of histone 3 (H3K4me1). Remarkably, both H3K27ac and H3K4me1 ChIP-seq signals mirrored the DNase-seq signal and were detected in particular at kidney and heart DHSs exhibiting liver-specific BMAL1 peaks (Fig. 2C,F; Supplemental Fig. S2D).

In summary, our data indicate that only a minority of BMAL1-binding sites are located at tissue-specific DHSs. For most tissue-specific BMAL1 peaks, chromatin is accessible in other tissues and harbors histone modifications that mark transcriptionally active DHSs, albeit to lower levels. Importantly, some BMAL1 DNA binding can be detected at those sites, although at levels too low to be categorized as ChIP-seq peaks by peak-calling algorithms.

Tissue-specific BMAL1 peaks are enriched for ts-TF DNA-binding motifs

Several mechanisms may contribute to tissue-specific BMAL1 binding. First, BMAL1 DNA binding may only
rely on the presence of its binding motif in an accessible DHS. Under this scenario, BMAL1-binding signal would correlate with DNase-seq signal, and differences between tissues would mostly depend on the proportion of cells harboring an accessible DHS and/or the extent to which the DHS is accessible. Alternatively, tissue-specific BMAL1 binding may be, as suggested for some other TFs (Hu and Gallo 2010; Gertz et al. 2013; Meireles-Filho et al. 2014), facilitated by ts-TFs that bind at sites nearby BMAL1. Here, BMAL1 DNA-binding signal would not correlate well with DNase-seq signal, and motifs for ts-TFs would be enriched at tissue-specific BMAL1 peaks.

To assess whether either or both of these mechanisms promote tissue-specific BMAL1 binding, we first compared BMAL1 ChIP-seq signal with DNase-seq signal at liver-, kidney-, and heart-specific BMAL1 peaks and found that both signals were correlated in the heart but not in the liver and kidney (Fig. 3A). Differences in BMAL1 ChIP-seq signal of >16-fold are detected at liver DHSs displaying similar DNase-seq signals, and similar BMAL1 ChIP-seq signals were observed at DHSs harboring differences of DNase-seq signal of >30-fold in both the liver and kidney (Fig. 3A). Importantly, these differences are independent of the number of E-boxes within each DHS (Fig. 3B, Supplemental Fig. S3A). These results thus indicate that, at least for the liver and kidney, the extent to which the chromatin is accessible cannot solely explain the differences in tissue-specific BMAL1 binding.

To determine whether ts-TFs may be involved, we performed a motif enrichment analysis at tissue-specific BMAL1 peaks using HOMER. As expected, E-boxes are enriched at tissue-specific BMAL1 peaks for all three tissues, along with a few additional motifs for ubiquitously expressed TFs (u-TFs) (Fig. 3C,D, Supplemental Table S2). However, the majority of the motifs were significantly enriched in only one or two tissues, with many of them corresponding to binding sites of well-known ts-TFs like Foxa1 and Mef2a/b (Fig. 3C,D). To verify that these TFs are legitimate ts-TFs, we analyzed their levels of expression in the liver, kidney, and heart using public mouse RNA-seq data sets (Zhang et al. 2014) and the Genotype Tissue Expression (GTex) portal for human tissues (The GTEx Consortium 2013). For almost all of the >20 TFs that we examined, mRNA levels were specific to one or two tissues, confirming that these TFs are true ts-TFs (Fig 3E; Supplemental Fig. S3B,C). Importantly, the tissue-specific pattern of expression matched particularly well with the tissue specificity of the motif enrichment, suggesting that these TFs bind DNA in a tissue-specific manner.

Genomic footprints for ts-TFs are specifically enriched at tissue-specific BMAL1 peaks

The concordance between ts-TF motif enrichment and expression pattern (Fig. 3C–E; Supplemental Fig. S3B,C) does not directly show that ts-TFs bind at BMAL1 DHSs in a tissue-specific manner. To further investigate this possibility, we took advantage of the genomic footprints left by chromatin-associated proteins following treatment with DNase I (Hesselberth et al. 2009). This analysis, which has been used successfully to define the genome-wide DNA-binding profiles of dozens of TFs (Neph et al.
Figure 3. Motifs and footprints for ts-TFs are enriched at tissue-specific BMAL1 enhancers. (A) Correlation between DNase-seq and BMAL1 ChIP-seq signals (calculated at BMAL1 peak center ±250 bp) at liver-, kidney-, and heart-specific BMAL1 peaks. (B) Correlation between DNase-seq and BMAL1 ChIP-seq signal for liver-specific BMAL1 peaks harboring one E-box only. (C) Enrichment of TF motifs at liver-, kidney-, and heart-specific BMAL1 peaks, performed using HOMER at BMAL1 peak center ±100 bp. Enrichments are shown if q-value was <0.05. (D) Fold-enrichment of TF motif over background at tissue-specific BMAL1 peaks (BMAL1 peak center ±100 bp; q-value < 0.05). (E) mRNA expression in the mouse liver, kidney, and heart of Bmal1 and TFs whose motifs were enriched at BMAL1 ChIP-seq peaks. (RNA-seq data sets from Zhang et al. 2014). Groups with different letters are significantly different. P-value < 0.05, one-way ANOVA. (F) Distribution of DNase I footprints identified using pyDNase and detected at BMAL1 peak center ±100 bp, and parsed based on the tissues in which they were found. (G) Motif enrichment of TFs performed at DNase I footprints identified in liver-, kidney-, and heart-specific BMAL1 peaks (footprint center ±15 bp). Enrichments are displayed if q-value was <0.05, and colored in gray if no significant footprint is detected in any of the three tissues. (H) Heat map representing DNase I cuts at BMAL1 peaks containing an E-box and an HNF6 motif (top) or a CEBP (bottom). DNase I cut signal is centered on the E-box and sorted based on the distance between the E-box and the liver-specific TF motif. (I) Quantification of DNase I cut signal at BMAL1 peaks containing an E-box and an HNF6 motif (top) or a CEBP (bottom) in the liver of wild-type (left) or Bmal1−/− (right) mice. Quantification was performed using signal centered on E-boxes (left) or on the ts-TF motif (middle). As control, quantification was also performed at HNF6 or CEBPA peaks that do not harbor a BMAL1 ChIP-seq peak (right; signal corresponds to the average of 1000 decile-normalized iterations).
This analysis revealed that most footprints found at tissue-specific BMAL1 peaks are significantly enriched in the corresponding tissue. For example, 60% of the footprints found at liver-specific and heart-specific BMAL1 peaks are only detected in the liver and heart, respectively [Fig. 3F; Supplemental Fig. S3F]. Conversely, a large fraction of the footprints identified at common BMAL1 peaks are detected in all three tissues [Fig. 3F; Supplemental Fig. S3F]. To identify the proteins generating these footprints, we performed a motif analysis restricted to the footprint genomic locations found at liver-, kidney- and heart-specific BMAL1 peaks. While motifs for CLOCK:BMAL1 and some u-TFs are enriched in all three tissues, most significantly enriched motifs correspond to ts-TFs binding sites [Fig. 3G; Supplemental Fig. S3G; Supplemental Table S3]. For example, footprints are enriched at motifs for Hnf6, Foxa1, Cebp, Hnf1, and Hnf4a in the mouse liver, and for Hnf1, Hnf4a, and Tead in the kidney. We noticed that several ts-TFs whose motifs are enriched at BMAL1 DHSs [Fig. 3C] do not exhibit a significant footprint [e.g., Foxo3, Srebp, Pax8, Gata4, and Ets1] [Fig. 3G], suggesting that they do not bind to BMAL1 DHSs despite the presence of a cognate motif. However, the absence of footprints may also reflect the inability of these TFs to generate distinctive footprints because of DNA residence time and DNase I cut bias, as shown previously for a few other TFs [He et al. 2014; Sung et al. 2014].

Using heat maps, we then examined DNase I cuts at BMAL1 peaks and found that DHSs exhibit footprints at E-box and motifs for ts-TFs, suggesting that ts-TFs may be bound to DNA concurrently with CLOCK:BMAL1 and some u-TFs are enriched in all three tissues, most significantly enriched motifs correspond to ts-TFs binding sites [Fig. 3G; Supplemental Fig. S3G; Supplemental Table S3]. For example, footprints are enriched at motifs for Hnf6, Foxa1, Cebp, Hnf1, and Hnf4a in the mouse liver, and for Hnf1, Hnf4a, and Tead in the kidney. We noticed that several ts-TFs whose motifs are enriched at BMAL1 DHSs [Fig. 3C] do not exhibit a significant footprint [e.g., Foxo3, Srebp, Pax8, Gata4, and Ets1] [Fig. 3G], suggesting that they do not bind to BMAL1 DHSs despite the presence of a cognate motif. However, the absence of footprints may also reflect the inability of these TFs to generate distinctive footprints because of DNA residence time and DNase I cut bias, as shown previously for a few other TFs [He et al. 2014; Sung et al. 2014].

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Moreover, dual E-boxes—i.e., two E-box motifs separated by 6 or 7 nucleotides and which have been proposed to be a preferred CLOCK:BMAL1-binding site (Paquet et al. 2008)—are also more enriched at common BMAL1 peaks (Fig. 4F). Characterization of the location of BMAL1 peaks revealed that peaks common to all three tissues were predominantly located at promoter/transcription start site (TSS), whereas tissue-specific BMAL1 peaks were more consistently located within introns and intergenic regions (Fig. 4G). Finally, nucleosome signal at common BMAL1 peaks was lower compared with other groups (Fig. 4H; Supplemental Fig. S4H). This result, which is likely due to differences in BMAL1 peak location between groups, suggests that DNA is more accessible at peaks common to all three tissues.

Together, these results indicate that BMAL1 peaks common to all three tissues, which target genes that are involved in generic biological processes, are enriched for E-boxes, are predominantly localized at promoter regions/TSS, and exhibit less nucleosome signal. These features likely contribute to the higher BMAL1 ChIP-seq signal observed at those sites.

BMAL1 DNA binding contributes only partially to rhythmic gene expression

To determine whether tissue-specific BMAL1 binding contributes to generating tissue-specific circadian transcription, we analyzed BMAL1 target gene expression in the mouse liver, kidney, and heart using public data sets that assessed the rhythmic transcriptome in those tissues (Zhang et al. 2014). In agreement with the literature (Rey et al. 2011; Menet et al. 2012), genes targeted by BMAL1 exhibit a higher proportion of rhythmic mRNA expression than those not targeted by BMAL1. Specifically, rhythmic expression of BMAL1 targets in the liver, kidney, and heart is increased by approximately twofold compared with all genes and by ~1.3-fold to 1.5-fold compared with other groups (Fig. 4E). Moreover, dual E-boxes—i.e., two E-box motifs separated by 6 or 7 nucleotides and which have been proposed to be a preferred CLOCK:BMAL1-binding site (Paquet et al. 2008)—are also more enriched at common BMAL1 peaks (Fig. 4F). Characterization of the location of BMAL1 peaks revealed that peaks common to all three tissues were predominantly located at promoter/transcription start site (TSS), whereas tissue-specific BMAL1 peaks were more consistently located within introns and intergenic regions (Fig. 4G). Finally, nucleosome signal at common BMAL1 peaks was lower compared with other groups (Fig. 4H; Supplemental Fig. S4H). This result, which is likely due to differences in BMAL1 peak location between groups, suggests that DNA is more accessible at peaks common to all three tissues.

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with genes targeted by BMAL1 in other tissues (Fig. 5A).

Surprisingly, genes targeted by BMAL1 in all three tissues (group 1) do not exhibit substantially more rhythmic expression despite increased BMAL1 ChIP-seq signal, increased number of E-boxes, and preferential peak location at promoter/TSS (Fig. 5A). Moreover, the majority of BMAL1 target genes are not rhythmically expressed in any of the three tissues (Fig. 5A). These results thus suggest that although BMAL1 DNA binding contributes to increased rhythmic expression, it is not sufficient to drive rhythmic transcription.

Consistent with this notion, BMAL1 targets common to all three tissues (group 1) display a widely heterogeneous rhythmic output, with most genes being rhythmic in only one tissue (83 genes out of 125 rhythmically expressed genes) (Fig. 5B). In fact, only 16 genes targeted by common BMAL1 peaks are rhythmically expressed in all three tissues, and most of them are core clock genes. We also found that many genes targeted by BMAL1 in one tissue were rhythmic only in another tissue (Fig. 5B). To characterize the mechanisms underlying this poor overlap between BMAL1 binding and rhythmic gene expression, we examined the profiles of DNase-seq and BMAL1 ChIP-seq data sets in the liver, kidney, and heart. While we were unable to detect a unified mechanism that may explain the variability in BMAL1 transcriptional output, we identified two important features that likely contribute to rhythmic transcription.

First, similar BMAL1 ChIP-seq signal between tissues can result in significantly different levels and/or rhythmicity of target gene expression (Fig. 5C; Supplemental Fig. S5A). This observation was made at genes targeted by a unique DHS, suggesting that factors other than just BMAL1 contribute to the differences in mRNA expression, where the genes Coq10b and Abcf1 (C) represent two examples for which the activity of BMAL1 DHS likely contributes to the differences in mRNA expression. Second, we found that rhythmicity of BMAL1 target genes was often associated with higher levels of expression and more DHSs, thus suggesting that rhythmic gene expression results from functional interactions between BMAL1 DHSs and other DHSs (Fig. 5D; Supplemental Fig. S5B). We also found a few instances where more DHSs were associated with decreased expression, suggesting that interactions between multiple DHSs may also

Figure 5. Transcriptional activities of BMAL1 DHS and other DHS contribute to BMAL1-mediated rhythmic transcription. [A] Percentage of rhythmically expressed gene for the seven groups of BMAL1-binding sites. Gene expression data originate from public microarray data sets (Zhang et al. 2014) and is considered rhythmic if q-value was <0.05 using JTK-cycle. [B] Overlap of rhythmic expression for peaks common to all three tissues (group 1), and for liver-, kidney-, and heart-specific BMAL1 peaks (groups 5, 6, and 7, respectively). The number of genes that are rhythmically expressed for the mouse liver (blue), kidney (green), and heart (red) is displayed, along with the number and percentage of nonrhythmically expressed genes (AR). [C, D] mRNA expression (left) and genome browser view (right) of BMAL1 ChIP-seq and DNase-seq signals in the mouse liver (blue), kidney (green), and heart (red). Rhythmic expression determined by JTK cycle is defined as three asterisks if q-value was <0.001, two asterisks if q-value was <0.01, and one asterisk if q-value was <0.05. The genes Coq10b and Abcf1 (C) represent two examples for which the activity of BMAL1 DHS likely contributes to the differences in mRNA expression, where the genes Dusp7 and Cont (D) represent two examples for which the activity of other DHS likely contributes to BMAL1-mediated rhythmic transcription.
result in decreased transcription activation (Supplemental Fig. S6C).

In summary, our results indicate that, as reported recently [Trott and Menet 2018], BMAL1 DNA binding is not sufficient to drive rhythmic transcription. They also suggest that rhythmic BMAL1 target gene expression likely results from [1] interactions between DHSs bound by BMAL1 and other DHSs and [2] the activity of each DHS, including those bound by BMAL1.

**BMAL1-bound DHSs physically interact with other DHSs**

To determine the extent to which BMAL1 target gene transcription relies on physical interactions between DHSs, we performed RNA Pol II chromatin interaction analysis by paired-end tag (ChIA-PET) sequencing in the mouse liver at ZT6 and ZT18 with three biological replicates per time point. By incorporating a Pol II ChIP step, this technique identifies interactions between enhancers and/or TSS of genes that are being transcribed [Fig. 6A; Li et al. 2012; Kieffer-Kwon et al. 2013]. To assess the efficiency of our mouse liver Pol II ChIA-PET in detecting chromatin interactions at transcriptionally active genes, we determined the number of PETs mapped to a single gene based on gene transcription using public mouse liver Nascent-seq (genome-wide sequencing of nascent RNA) data sets [Menet et al. 2012]. We found that most PETs mapped to transcriptionally active genes, with 23.4% of all PETs located within the top 10% of genes transcribed in the mouse liver [Fig. 6B; Supplemental Fig. S7A; Supplemental Table S4]. Analysis of PETs with reads located in two different DHSs and visualization of chromatin interactions at mouse liver BMAL1 target genes further confirmed that actively transcribed genes harbor many PETs mapping to different DHSs, including some bound by BMAL1 [Fig. 6C,D]. These data indicate that DHSs located within the same gene physically interact via gene looping, and suggest that transcriptional activation results from the synergistic interaction

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**Figure 6.** Analysis of chromatin interactions by RNA Pol II ChIA-PET in the mouse liver. (A) Illustration of the ChIA-PET technique. (B) Number of PETs with both reads mapped to the same gene and parsed based on gene nascent RNA expression in the mouse liver. Red circles represent the average PET number for each decile ±95% confidence intervals. (C) Average number of PETs per gene in the mouse liver with both reads located in different DHS and mapped to the same gene, parsed based on gene nascent RNA expression. Error bars represent the 95% confidence intervals. (D) Genome browser view of mouse liver BMAL1 ChIP-seq [this study], DNase-seq (ENCODE), and Pol II ChIP-seq [Sobel et al. 2017]. PETs with both reads mapped to DHS are in red, while PETs with one read mapped to a DHS are in orange and those not mapped to a DHS are in gray. (E–G) Percentage of PETs detected at ZT6 (white bar and dashed black line) or ZT18 (gray bar and solid black line), displayed as the average ± S.E.M. of three independent experiments based on the type of PETs, the rhythmicity of gene expression, and the phase of expression. (*) P < 0.05 between ZT6 and ZT18; (#) an interaction found in the two-way ANOVA [P < 0.05] between the phase of expression and the time of the ChIA-PET experiment (ZT6 or ZT18). Triangles represent DHSs and may be located within a gene or not.
between DHSs. To assay whether rhythmic gene expression is associated with changes in chromatin interactions, we determined the number of PETs between two DHSs based on the phase of rhythmic gene expression. We found small day versus night differences in chromatin interactions in the mouse liver, with ~10% more PETs between two DHSs at ZT6 for genes with a peak of expression at ZT9–15 [Fig. 6E]. In addition, analysis of interactions between a DHS and a non-DHS site, e.g., a site with elongating Pol II that is expected to better reflect transcriptional activity revealed that DHS–non-DHS interactions exhibit day versus night differences, with more interactions at ZT18 for genes expressed from ZT15 to ZT3 [Fig. 6F,G].

A few recent reports suggested that rhythmic interactions between BMAL1 DHSs and other DHSs underlie the rhythmic transcription of BMAL1 target genes (Aguilar-Arnal et al. 2013; Mermet et al. 2018; Yeung et al. 2018). To investigate this possibility, we quantified the number of PETs mapped to BMAL1 target genes. We found that at rhythmically expressed BMAL1 target genes the number of chromatin interactions between two DHSs containing at least one BMAL1 peak is higher at ZT6 when BMAL1 binding is maximal [Fig. 7A,B]. This higher number of PETs at ZT6 versus ZT18 was specifically observed for rhythmically expressed targets peaking at ZT3–ZT15 (i.e., at times corresponding relatively well with BMAL1 DNA binding) and after incorporation of chromatin interactions involving distal BMAL1-bound DHSs, which account for ~15%–20% of DHS–DHS interactions [Fig. 7A, B]. In addition, analysis of interactions between a BMAL1 peak and a non-DHS site revealed that while the proportion of PETs between ZT6 and ZT18 was similar for rhythmically and arrhythmically expressed targets, more PETs were found at ZT18 for the target genes rhythmically expressed at a similar phase [Fig. 7C,D].

To further characterize chromatin interactions between BMAL1-bound DHSs and other DHSs, we analyzed the type of DHSs with which BMAL1-bound DHSs interact (i.e., liver-specific DHS or DHS common to the kidney and/or heart) as well as their genomic location (i.e., promoter, intron, etc.). We found that interactions between BMAL1-bound DHSs and DHSs common to several tissues were enriched for rhythmically expressed genes [Fig. 7E], and that interactions with DHSs located in introns were enriched for rhythmically expressed genes with a peak of expression at ZT9–ZT15, i.e., a few hours after maximal BMAL1 DNA binding [Fig. 7F]. Future experiments will be required to determine whether these differences in the type and location of chromatin interactions are relevant for BMAL1-mediated rhythmic gene expression.

Taken together, these results indicate that BMAL1-bound DHSs interacts with other DHSs, and that these interactions are more prevalent when BMAL1 is bound to DNA at ZT6 for rhythmic targets expressed at ZT3–ZT15. Because the number of interactions is not rhythmic for arrhythmically expressed targets, this suggests that rhythmic chromatin interactions between BMAL1-bound DHSs and other DHS may contribute to rhythmic gene expression.

Discussion

Although the same clockwork mechanism is found in essentially all tissues, clock-controlled gene rhythmic expression is mainly tissue-specific, reflecting the large number of biological functions that are clock controlled [Panda et al. 2002; Storch et al. 2002; Zhang et al. 2014; Mure et al. 2018]. By characterizing BMAL1 cistromes in three mammalian tissues, we show that tissue-specific circadian transcriptional programs can be directly initiated by the core circadian clock. We provide evidence that BMAL1 DNA-binding profiles are largely tissue-specific, which coincides with tissue-specific chromatin accessibility and co-binding with ts-TFs. Our analysis of BMAL1 target gene expression also indicates that rhythmic CLOCK:BMAL1 transcriptional output relies, at least in part, on the rhythmic interaction between BMAL1-bound DHSs and other DHSs. Given that some of these other DHSs are either tissue-specific or more accessible in specific tissues [e.g., Fig. 5D; Supplemental Fig. S5B], our data suggest that enhancer–enhancer interactions can drive rhythmic BMAL1 target gene expression in a tissue-specific fashion [Fig. 7E]. Taken together, chromatin accessibility and enhancer–enhancer interactions may explain how the circadian clock generates tissue-specific circadian transcriptional programs, thereby regulating biological functions in a tissue-specific manner. We anticipate that these mechanisms apply to other TFs that regulate gene expression in a tissue-specific manner.

Recent characterization of the chromatin-accessible landscape revealed that most TFs bind to regions that are hypersensitive to nuclease digestion [Thurman et al. 2012; Yue et al. 2014]. Because many of these DHSs are tissue specific [Supplemental Fig. S2A], it is not surprising that differences in the openness of the chromatin between tissues can promote tissue-specific BMAL1 binding. However, the majority of tissue-specific BMAL1 peaks are located in chromatin regions accessible in other tissues [Fig. 2C; Supplemental Fig. S2C,D]. DHSs are clusters of TF-binding motifs, and TF recruitment to DHSs often results from cooperative binding between TFs. In particular, ts-TFs can cooperate with u-TFs to facilitate DNA binding: CEBPB cooperates with glucocorticoid receptor in the mouse liver [Grøntved et al. 2013], FOXA1 and GATA3 with the estrogen receptor in primary breast tumors [Hurtado et al. 2011; Theodorou et al. 2013], and OPA and SERPENT with Drosophila CLK:CYC [Mireles-Filho et al. 2014]. Our finding that binding motifs, genomic footprints, and ChIP-seq signal for liver-specific TFs are enriched at liver-specific BMAL1 peaks, but not at common peaks strongly suggests that this ts-TF/u-TF cooperation applies to BMAL1 and that ts-TFs facilitate tissue-specific CLOCK:BMAL1 binding [Fig. 7C].

By analyzing the genome-wide nucleosome positioning in the mouse liver over the course of the day, it has recently been shown that CLOCK:BMAL1 promotes rhythmic nucleosome removal at its binding sites and proposed that CLOCK:BMAL1 is a pioneer-like TF [Menet et al. 2014]. Our findings that BMAL1 binds in a tissue-specific manner even at DHSs common to several tissues seems
contradictory to BMAL1 pioneering-like function. Instead

of generating new DHSs (an expected feature from pioneer

TFs) (Zaret and Carroll 2011), our data rather suggest that

BMAL1 DNA binding prevents the folding of DNA around

histones and thus nucleosome formation. As a result,

BMAL1 DNA binding would increase the exposure of na-

ked DNA for other TFs and DNase I and facilitate their

binding on DNA. This scenario, which is consistent

with the analysis of nucleosome signal at tissue-specific

BMAL1 peaks in $\mbox{Bmal1}^{-/-}$ mice (Supplemental Fig.

S4H), would explain why DNase I signal is decreased in

$\mbox{Bmal1}^{-/-}$ mice around not only E-boxes but also ts-TFs

motifs (Fig. 3H,I). Conversely, ts-TFs may play a similar

role at tissue-specific DHSs, i.e., their binding to DNA may facilitate the access of CLOCK:BMAL1 to exposed E-boxes by preventing nucleosome formation. This mechanism, which has been proposed previously to illustrate cooperative TF binding to DHSs (Mirny 2010), may explain at least in part tissue-specific BMAL1 binding at DHSs common to several tissues. Future experiments will be required to experimentally validate this hypothesis and eventually clarify BMAL1 pioneering like function.

Genes targeted by BMAL1 in all three tissues do not ex-
hibit a higher rate of rhythmic expression and display a re-
markable heterogeneity in the rhythmic output (Fig. 5A,
B), indicating that even robust BMAL1 binding is not
sufficient to drive rhythmic transcription. As recently suggested [Trott and Menet 2018], this may be explained by the contribution of u-TFs in regulating the transcriptional activity of BMAL1 DHSs and promoting gene expression. However, our analysis suggests an additional level of regulation that includes DHSs unbound by BMAL1, and through which interactions between BMAL1-bound DHSs and other DHSs would contribute to the regulation of rhythmic gene expression (Fig. 5D). To characterize whether DHSs not bound by BMAL1 might contribute to rhythmic BMAL1 transcriptional output, we conducted mouse liver Pol II ChIA-PET to identify physical interactions between DHSs within genes engaged in active transcription (Fig. 6). In agreement with the literature, we found that the number of DHS–DHS interactions correlates with transcriptional levels. Relevant to how the circadian clock might regulate rhythmic gene expression, we identified a large number of interactions between BMAL1-bound DHSs and other DHSs, suggesting that BMAL1 target gene transcription relies more on the functional interaction between DHSs rather than just BMAL1-bound DHSs. In addition, and consistent with previous reports that used the 4C technique (Águilar-Arnal et al. 2013; Mermet et al. 2018; Yeung et al. 2018), we found that the number of interactions between BMAL1 DHSs and other DHSs is higher at ZT6; i.e., at the time of maximal BMAL1 DNA binding. Importantly, this higher number of chromatin interactions at ZT6 was observed for rhythmically expressed target genes that peak during the day and not at arrhythmically expressed targets, suggesting that BMAL1-mediated rhythmic transcription relies on the capacity of BMAL1 to regulate rhythmic enhancer–enhancer interactions. These results are concordant with recent reports showing that rhythmic transcription in mammals is associated with rhythmic long-range interactions between cis-regulatory elements and rhythmic gene looping [Águilar-Arnal et al. 2013; Kim et al. 2018; Mermet et al. 2018; Yeung et al. 2018]. While the mechanisms underlying rhythmic enhancer–enhancer interactions remain vastly unknown, they likely involve the Mediator complex and transcriptional regulators recognizing histone modifications associated with transcriptional activation [Kim et al. 2018]. In addition, our and others’ findings raise the hypothesis that the expression of BMAL1 target genes relies on the relative contribution of BMAL1-bound DHSs versus other DHSs, and that changes in the activity of other DHSs can impair the rhythmic expression of BMAL1 targets without affecting the molecular clockwork mechanism. Such a mechanism may well explain why circadian transcriptional programs can be reprogrammed by environmental challenges without affecting much of the clockwork mechanism [Eckelman et al. 2013; Murakami et al. 2016; Sato et al. 2017].

In summary, our results provide novel insights into how BMAL1 regulates rhythmic gene expression in a tissue-specific manner and shed light on the role of enhancer–enhancer interactions in generating circadian transcriptional programs. We anticipate that these findings will be relevant for our understanding of how the circadian clock regulates a wide array of biological functions under normal and diseased states and will apply more generally to how other TFs regulate tissue-specific gene expression.

Materials and methods

Animals

Male C57BL/6j and Bmal1−/− mice were housed under 12 h light:12 h dark (LD12:12) with food and water available ad libitum. Bmal1−/− were kindly provided by Christopher Bradfield (Bunger et al. 2000). The age of animals collected was between 3 and 6 mo old. All experiments were approved by the Texas A&M University Institutional Animal Care and Use Committee.

BMAL1 ChIP-seq

BMAL1 ChIP-seq were performed as described previously [Menet et al. 2014] with modifications. Formaldehyde crosslinked nuclei were purified by centrifugation on a 24% sucrose cushion at 20,000g, washed, and sonicated to obtain chromatin fragments of about 100–600 base pairs (bp) in length. Immunoprecipitation was performed using 1 µL of BMAL1 antibody [chicken anti-BMAL1] and BMAL1 ChIP efficiency was verified by qPCR. ChIP-seq libraries (n=3 mice per tissue) were generated using NEBNext ChIP-seq library preparation master mix set as per the manufacturer’s instructions and sequenced using an Illumina NextSeq.

Sequencing data sets and alignment to the mouse genome

All public data sets were downloaded as fastq or Short Read Archive (SRA) file formats (accession numbers are provided in the Supplemental Material). To avoid issues due to the utilization of different protocols/procedures for each tissue, each of the liver, kidney, and heart ChIP-seq, DNase-seq and mRNA expression data sets used in cross-comparisons were generated from the same research laboratory. ChIP-seq and DNase-seq data sets were aligned to the mouse genome (version mm10) using bowtie2 with the parameters: -x and --end-to-end. Uniquely mapped reads were only considered for analysis, and up to three duplicated sequences were kept for each BMAL1 ChIP-seq data set. Visualization files were generated using bedtools and normalized to 10,000,000 reads. Input files were processed individually and then merged as bam files using samtools.

For DNase-seq data sets, bam files from all technical and biological replicates were merged and no downsampling was performed. Reads from the RNA-seq data set were trimmed using fastx_trimmer with the following parameters -f 1 -l 100 -Q 33, and aligned to the genome using STAR with the default parameters and the option: -outFilterIntronMotifs RemoveNoncanonical. Gene expression data were retrieved using cufflinks and the genome version GRCh38.p5_M14 and default parameters.

Sequencing data set analysis

Peak calling for both BMAL1 ChIP-seq and DNase-seq data was performed with findPeaks from the HOMER suite. A minimum local enrichment of fourfold was set up as necessary, and the following parameters were used: -style factor and -i [for BMAL1 ChIP-seq] or -style dnas and -region [for DNase-seq]. Overlap between ChIP-seq or DNase-seq peaks was determined using the function intersectBed from Bedtools suite with default parameters. Heat maps were using the Rscript pheatmap.R.

BMAL1 peaks were assigned to their target genes using the Perl script annotatePeaks.pl from the HOMER suite. The HOMER
gene annotation script outputs each peak into the following categories: [1] promoter–TSS, corresponding to TSS −10 kb to TSS +1 kb, [2] transcription termination site (TTS), corresponding to TTS −100 bp to TSS +1 kb, [3] exons, [4] introns, and [5] intergenic, which corresponds to peaks located upstream of the TSS by >10 kb and downstream from the TTS by >1 kb. Intron and exon assignments were not adjusted from the output of HOMER annotatePeaks.pl. BMAL1 peaks labeled as intergenic were not assigned to a target gene.

ChIP-seq and DNase-seq signal was calculated using scripts from Bedtools and the uniquely mapped reads and the genomic coordinates of BMAL1 ChIP-seq peaks [Supplemental Table S1] or mouse liver DHS peaks [Supplemental Table S5]. Signal was calculated in a ±250-bp region from the peak center for DNase-seq as well as for BMAL1 and other TF ChIP-seq signal and in a ±1-kb region from the peak center for H3K27ac ChIP-seq signal.

Detection of footprints was performed using the python script wellington_footprints.py from the pyDNase suite. All parameters were set to default, and a P-value of ~0.20 was used along with a false discovery rate of 0.01. Motif analysis was performed at BMAL1 DNA-binding sites using the perl script findMotifsGenome.pl from the HOMER suite using the parameter -size 200. Motifs were considered as significantly enriched if the q-value was <0.05. Motif enrichment was calculated based on the background from the output of findMotifsGenome.pl. The genomic location of DNase I cuts was retrieved from DNase-seq data sets by reporting the position and strandness of the first nucleotide of each read in a bam file. Average DNase I cut signal was calculated at E-boxes and other TF-binding motifs using scripts from Bedtools. Quantification of the average DNase I cut signal at control peaks was performed by randomly selecting an equivalent number of TF ChIP-seq peaks without BMAL1 to the number of peaks with BMAL1. Because most TF ChIP-seq peaks with BMAL1 signal exhibit strong DNase-seq signal (i.e., most of them are within the strongest DHSs), the random selection was performed by matching the number of peaks within each DNase-seq signal decile. This process was repeated 1000 times, and the average of 1000 iterations calculated.

Generation of footprints for E-boxes and dual E-boxes was performed with the perl script seq2profile.pl from the HOMER suite. The E-boxes considered for analysis were CACGTG, CACNGN, and CACGTT, and the dual E-box motif tolerated up to two mismatches between the two E-boxes and contained a spacer of either six or seven base pairs. Gene ontology was performed using the perl script annotatePeaks.pl from the HOMER suite with the parameter -go and mm10 genome.

Rhythmic expression of BMAL1 target genes was determined using public microarray data sets performed in the same research laboratories [Zhang et al. 2014]. Files containing expression values for each microarray probe were downloaded from the NCBI Web site [GSE54652], and no analysis of the original files was performed. Rhythmic gene expression was determined using JTKsite (GSE54652), and no analysis of the original files was performed in the same research laboratories using public microarray data sets performed in the same research laboratories the parameter -go and mm10 genome.

The genomic location of DNase I cuts was retrieved from DNase-seq data sets by reporting the position and strandness of the first nucleotide of each read in a bam file. Average DNase I cut signal was calculated at E-boxes and other TF-binding motifs using scripts from Bedtools. Quantification of the average DNase I cut signal at control peaks was performed by randomly selecting an equivalent number of TF ChIP-seq peaks without BMAL1 to the number of peaks with BMAL1. Because most TF ChIP-seq peaks with BMAL1 signal exhibit strong DNase-seq signal (i.e., most of them are within the strongest DHSs), the random selection was performed by matching the number of peaks within each DNase-seq signal decile. This process was repeated 1000 times, and the average of 1000 iterations calculated.

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Graphs in Supplementary Fig. S3C displaying the RPKM (reads per kilobase per million mapped reads) values of different TFs in human livers, kidneys, and hearts (atrial appendage and left ventricle) were retrieved from GTEx portal in April 2016.

Mouse liver Pol II ChIA-PET

Mouse liver Pol II ChIA-PET experiments were performed at ZT6 and ZT18 with three biological replicates according to published protocols [Li et al. 2010] with some modifications. Liver nuclei were either dual cross-linked with 1.5 mM EGS and 1% formaldehyde (two experiments) or single cross-linked with 1% formaldehyde (one experiment). Sonicated chromatin was immunoprecipitated with an anti-RNA Pol II 5WG16 antibody. ChIA-PET libraries were sequenced on a HiSeq 2000 or a MiSeq. Additional information is in the Supplemental Material.

Sequencing and computational analysis of Pol II ChIA-PET libraries

ChIA-PET reads from the fastq files were processed to extract the tags 1 and 2 along with their accompanying half-linker code using a custom-made Python script and to generate two fastq files (R1 and R2 files) containing the sequence identifier, the raw sequence, and the sequence quality values for each tag. These two files were then aligned to the mouse genome (mm10 version) as paired-end reads using bowtie2. Only paired tags with both reads mapping uniquely to the mouse genome were considered in our analysis. PETs were parsed based on the half-linker barcodes into nonchimeric PETs (specific products) or chimeric PETs (nonspecific products) and duplicated PETs were removed for both chimeric and nonchimeric products. PETs with a tag location shifted by 1 bp compared with an existing PET were also considered as PCR duplicates and removed.

PETs with both reads on the same chromosome and with a distance between reads ≥500 bp were only considered in our analysis. For all PETs, each of the two tags was extended to 200 bp (tag location ±100 bp) and this tag genomic location (chr:start-end) was used to map tags to a gene, and a DHS, using intersectBed from bedtools. Mapping to DHS was performed using a more stringent analysis of the mouse liver DNase-seq data sets from ENCODE [Supplemental Table S5] and which mostly reports stronger DHSs (see Fig. 6D). DHSs harboring a mouse liver BMAL1 peak were identified with intersectBed between the DHS peak list described above and the list of mouse liver ChIP-seq peaks generated in this manuscript [Supplemental Table S1]. To validate that mouse liver PETs contribute to gene transcription [Fig. 6B,C], we used public mouse liver Nascent-seq data sets, and averaged values for each of the 12 independent samples. Finally, we considered genes to be rhythmically expressed based on the analysis of the microarray data sets from Zhang et al. [2014] as described above.

Statistical analysis

Statistical analysis was carried out in JMP version 12.0.1. ChIP-seq and DNase-seq signals were analyzed using a Kruskal-Wallis test and post-hoc analysis with a Wilcoxon each pair test. Analysis of TF mRNA expression between the three tissues was performed using a one-way ANOVA. Analysis of the differences in BMAL1 peaks genomic locations was performed using a χ2 test, and differences in the number of BMAL1 peaks per genomic locations we analyzed by a Fisher’s exact test. Spearman correlation was used to determine the degree of correlation between signals (e.g., ChIP-seq with DNA-seq) or signal between tissues. Results were considered significant if P-value was <0.05 for the Kruskal-Wallis, Student’s t-test, and ANOVA tests, and P-value was <0.01 for Fisher’s exact test.
Data availability

The sequencing data sets generated in this study (BMAL1 ChIP-seq and Pol II ChIA-PET) have been deposited to Gene Expression Omnibus under the accession code GSE110604.

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Author contributions: J.R.B. and J.S.M. conceived and designed the research. J.R.B. performed most of the experiments and the bioinformatics analysis. J.S.M. performed the ChiA-PET experiments with the help of A.J.T. and J.R.B. B.G., C.A.O., and J.S.M. helped with the bioinformatics analysis. J.S. and H.V. performed the research. J.R.B. performed most of the experiments and the bioinformatics analysis. J.R.B. and J.S.M. wrote the manuscript.

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