A day in the life of chromatin: how enhancer–promoter loops shape daily behavior

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Each spring, we get out of bed 1 h ahead of our biological wake-up time due to the misalignment of internal clocks with the light–dark cycle. Genetic discoveries revealed that clock genes encode transcription factors that are expressed throughout many tissues, yet a gap has remained in understanding the temporal dynamics of transcription. Two groups now apply circular chromosome conformation capture and high-throughput sequencing to dissect how “time of day”-dependent changes in chromatin drive core clock oscillations. A surprise is the finding that disruption of enhancer–promoter contacts within chromatin leads to an advance in the “wake-up” time of mice. Furthermore, the assembly of transcriptionally active domains of chromatin requires the ordered recruitment of core clock transcription factors each day. These studies show that waking up involves highly dynamic changes in the three-dimensional positioning of genes within the cell.

Circadian clock genes, among the first ones identified to control behavior, are encoded by an autoregulatory feedback loop in which transcription factors in the forward limb activate their own repressors in the negative limb, generating oscillations with a periodicity of ∼24 h (Hardin et al. 1990). The discovery that the core clock is also widely expressed outside of the central nervous system in animals led to recognition that cell-intrinsic clocks in turn drive the oscillation of a vast number of genes, enabling anticipation of daily changes in the external environment imposed by the light–dark cycle. Circadian processes are also subjected to post-transcriptional and post-translational regulation, although, at its core, gene regulatory mechanisms remain central to understanding what makes a clock “tick.” Genomic approaches have identified distinct phases in clock transcription factor binding, histone modifications, and RNA polymerase II recruitment to DNA over the 24-h time scale (Koike et al. 2012), with a surprise being that rhythmic gene regulation occurs within enhancer regions that are far away in linear sequence from the proximate transcription start site of oscillating targets (Fang et al. 2014). A question has therefore arisen: How might interactions between regulatory enhancers and oscillating genes contribute to anticipatory behavior and physiology? To address this challenge, both the work in this issue of Genes & Development by Naef and colleagues (Mermet et al. 2018) and studies by Lazar and colleagues (Kim et al. 2018) exploit newly available tools from biochemistry and genomics to probe how chromatin reconfiguration evolves over the course of just 1 d. Especially surprising is the finding by Mermet et al. (2018) that genetic abrogation of a specific intronic enhancer element discovered through conformational analyses of chromatin alters rhythmic locomotor activity, suggesting that such loops are necessary for the maintenance of a behavior. Likewise, Kim et al. (2018) provide an unbiased survey of enhancer–promoter interactions, pinpointing a role of the repressor REV-ERβ as a regulator of the clock repressor Cryptochrome 1 (CRY1) and highlighting its function as a core component of the clock.

Just as discovery of the clock provided evidence for temporal control of gene transcription, insight into spatial control across the genome has advanced with the advent of next-generation sequencing. To this end, both the Naef group (Mermet et al. 2018) and the Lazar group (Kim et al. 2018) apply variations of chromatin analyses based on “3C” [chromatin conformation capture] approaches [Fig. 1]. The basic strategy is to cross-link chromatin-bound DNA and then identify the pairing of distant regulatory sequences through sequential steps of restriction endonuclease digestion followed by ligation, reverse cross-linking, and sequencing of the ligated fragments [for in-depth discussion, see Denker and de Laat 2016]. The key to this powerful technology is the concept that nucleotides within enhancers widely separated in linear sequence from the promoters that they target may occupy proximate spatial topologies due to enhancer–promoter looping. Such enhancer–promoter looping can be monitored by the paired proximate interactions (PPI) approach, which allows for visualization of the loops in 3D. First, chromatin is cross-linked, then digested with restriction enzymes, and then ligation of the DNA fragments is carried out. The ligated DNA is then sequenced and analyzed to identify looping events. This approach has been successfully applied to study the looping of enhancers in the context of the genome, allowing for the visualization of the spatial arrangement of enhancers relative to their target genes. The results of these studies have provided insights into the mechanisms by which enhancers control gene expression during development and disease.

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be frozen upon cross-linking (because, by definition, they are in proximity in three-dimensional space), thereby enabling subsequent ligation and sequence-based identification of topologically colocalized regulatory sequences.

Mermet et al. [2018] used circular chromosome conformation capture (4C), in which a target regulatory region is used as the “bait,” ligated contacts are circularized, and subsequent identification of bait contacts is performed by sequencing. One such bait used here was an established intragenic regulatory sequence within the gene encoding the clock repressor CRY1, previously identified in elegant work by Ukai-Tadenuma et al. [2011] as a modulator of rhythmic circadian oscillation. Contacts with the Cry1 intragenic enhancer from nearby regions peaked during the subjective night (near the zenith of Cry1 expression) and were diminished during the subjective day (the Cry1 nadir). Of note, these patterns were present in both the kidney and the liver, a finding consistent with the observation that enhancer regulation of core clock loci is conserved across tissues [Perelis et al. 2015]. In contrast, rhythmic intragenic promoter interactions were observed only in the liver for the gene encoding glycogen synthase 2, a clock output gene involved in glucose production, suggesting tissue specificity. Consistent with a role of the core circadian mechanism in driving Cry1 enhancer–promoter looping, 4C-seq (4C combined with sequencing) analyses in Bmal1 knockout mice displayed static elevated levels of Cry1. Since Bmal1 mutant animals are also deficient in the circadian repressors REV-ERBα and REV-ERBβ and have elevated levels of the activator RORγ, one possibility may be that the loss of repression leads to static chromatin assembly. Enhancer–promoter loops thereby establish a transcriptional feedback loop!

Following the concept that functional experiments are necessary to discern the meaning of specific interactions captured by 4C-seq, Mermet et al. [2018] generated a transgenic mouse harboring a 300-base-pair deletion within the intronic Cry1 enhancer. Animals bred to homozygosity with the loss of this single Cry1 enhancer display significantly shortened period length under constant conditions, a hallmark of circadian clock disruption and consistent with the shortening observed in Cry1 nullizygous mice, demonstrating the necessity of feedback repression in the core clock [Sato et al. 2006]. As predicted, rhythmic chromatin contacts between the enhancer and the promoter were also lost in the enhancer mutant mice. At the cellular level, single-molecule RNA fluorescence in situ hybridization (smRNA-FISH) further corroborated a requirement for enhancer–promoter looping in de novo transcription. Collectively, the findings of Mermet et al. [2018] begin to uncover a new level of molding in the emergence of an intact rhythmic organism. The aforementioned work by Mermet et al. [2018] in intact animals also resonates with previous cell-based assays using 4C followed by microarray chip hybridization to identify distant regions interacting with an intragenic regulatory motif within the gene encoding the D-albumin-binding protein (DBP), a clock output factor [Aguilar-Arnal et al. 2013].

Technologies for detecting long-range chromatin interactions continue to evolve, enabling exploration of genome topology across all possible regulatory enhancer–promoter loops. In the “all-to-all” approach, chromosome capture followed by high-throughput sequencing (Hi-C) allows for the generation of unbiased contact maps. With sufficiently deep sequencing (as performed by Kim et al. 2018), topologically associated domains (TADs) and smaller localized sub-TADs give unbiased information on proximal DNA ligations and the frequency of such contacts. This provides a way in which cis and trans chromosome interactions are quantified at a resolution sufficient for mapping individual sites of exon/intron, enhancer, and promoter contacts. Using Hi-C, Lazar and colleagues [Kim et al. 2018] analyzed the mechanisms governing enhancer–promoter looping in proximate interactions under ~300 kb. This “all-to-all” approach enabled unbiased identification of “time of day”-dependent interactions, with more observed during the subjective daytime, closer to the zenith of transcriptional activity of forward limb clock activators. Here, the studies also revealed correspondence between rhythmic chromatin interactions and rhythmic transcription, suggesting that topological transitions characterize rhythmic control of clock-controlled gene outputs. Intriguingly and echoing...
the findings of Mermet et al. [2018]. Cry1 enhancer-promoter looping was localized to a “time of day”-dependent region that emerged primarily during the subjective night. During the daytime, when expression of Cry1 is at its nadir, directed analysis further confirmed REV-ERBa binding to the Cry1 regulatory motif in an anti-phasic pattern with maximal expression of the gene, and liver-specific overexpression of REV-ERBa was sufficient to abrogate rhythmic Cry1 expression and enhancer-promoter looping. Genetic analyses of enhancer-promoter looping in the absence of REV-ERBa implicate a direct role for this factor in both core clock and clock output regulation. Indeed, the activity of REV-ERBa in modulating enhancer-promoter looping appears to involve the recruitment of corepressors containing nuclear receptor corepressor and histone deacetylase 3. These modulate histone 27 Lys27 acetylation and corresponding docking of the transcriptional elongation factor bromodomain-containing protein 4 (BRD4) and the looping factor Mediator complex 1 subunit, leading to the induction of RNA polymerase II. Collectively, the work of Kim et al. [2018] places REV-ERBa as an integral component of core circadian gene regulation. The techniques used here pave the way for future functional genomic studies to address how each component of the core clock and clock-driven processes mediate circadian dynamics and the diverse cellular physiologic events each day.

Chromatin conformation analyses now provide a new view of core clock regulation and highlight gaps in our understanding of how the circadian system prepares organisms for both anticipated and unexpected changes in the environment. Importantly, circadian transcriptional and epigenetic signatures are shared across tissues, yet unique tissue-specific programs are required for organismal homeostasis [Perelis et al. 2015]. Application of chromatin analyses to examine the extent to which core clock transcription factors and/or clock-recruited factors establish genomic topologies at the right time of day to impact physiology still remains an open area. Furthermore, mounting evidence suggests that core circadian processes exhibit dynamic responses to changes in the environment, raising the intriguing possibility that environmental signals might alter circadian chromatin programming. It appears that a new geometry is emerging, through which we can now view the shape of time.

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