MicroRNAs: a potential interface between the circadian clock and human health

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Abstract
The biochemical activity of a stunning diversity of cell types and organ systems is shaped by a 24-hour (circadian) clock. This rhythmic drive to a good deal of the transcriptome (up to 15% of all coding genes) imparts circadian modulation over a wide range of physiological and behavioral processes (from cell division to cognition). Further, dysregulation of the clock has been implicated in the pathogenesis of a large and diverse array of disorders, such as hypertension, cancer and depression. Indeed, the possibility of utilizing therapeutic approaches that target clock physiology (that is, chronotherapy) has gained broad interest. However, a deeper understanding of the underlying molecular mechanisms that modulate the clock, and give rise to organ-specific clock transcriptomes, will be required to fully realize the power of chronotherapies. Recently, microRNAs have emerged as significant players in circadian clock timing, thus raising the possibility that clock-controlled microRNAs could contribute to disorders of the human circadian timing system. Here, we highlight recent work revealing a key role for microRNAs in clock physiology, and discuss potential approaches to unlocking their utility as effectors of circadian physiology and pathophysiology.

Introduction
Deep within the brain ticks a 24-hour clock that imparts circadian control over virtually every aspect of human physiology. This clock is located in the suprachiasmatic nucleus (SCN), which is a tiny region of the ventral hypothalamus that contains approximately 20,000 neurons [1,2]. The inherent timing capacity of the SCN is derived from autonomous neuronal oscillators, which form a coherent and precisely phased pattern of rhythmic neuronal activity and hormonal output that serves as a phasing cue to ancillary oscillators found throughout the brain and peripheral organs [3,4]. The need for a circadian control center likely evolved as an adaptive response to the daily changes in the light-dark cycle resulting from the axial rotation of the earth. As such, the inherent clock timing process is tightly regulated by changes in the daily lighting cycle.

As discussed in detail below, dysregulation of the clock appears to play a central role in the genesis and progression of a broad range of disorders, and therapeutic approaches designed to target clock physiology (that is, chronotherapeutics) have been effective in treating a range of ailments, including seasonal affective disorder, asthma, hypertension and cancer [5-9]. Many of these treatment strategies are based on the integration of clock phase into the drug delivery schedule. However, if we are to develop novel chronotherapeutic approaches, then a deeper understanding of the underlying molecular mechanisms that modulate the clock, and give rise to organ-specific clock transcriptomes, will be required. To this end, recent work by a number of groups has revealed a role for microRNAs in clock physiology. MicroRNAs are small (approximately 22 nucleotides), single-stranded, non-coding RNA species that act as potent gene silencers and are important to a diverse array of physiological and pathophysiological processes [10-13]. As we will discuss in detail below, microRNAs appear to play distinct roles in clock physiology, and dysregulation of microRNA expression can lead to marked alterations in circadian timing and output. Hence, microRNAs present novel therapeutic targets for disorders of the circadian clock. In this review, we will provide an overview of circadian biology, and highlight genome-scale and integrated approaches designed to reveal the role of microRNAs in clock physiology. Finally, we will provide some perspective on microRNAs as potential therapeutic targets for circadian disorders. By understanding the role of microRNAs in circadian biology on molecular, cellular and organ-systems levels, a cohesive and integrated approach to chronotherapeutics can be developed.
Circadian timing
At a molecular level, the inherent 24-hour time-keeping capacity of the SCN stems from a well-characterized transcriptional/post-translation feedback loop (Figure 1). Central to this molecular clock is the rhythmic expression of the period (per1 and per2) and cryptochrome (cry1 and cry2) families of genes. Circadian transcription of per and cry genes is mediated by an E-box-binding basic helix-loop-helix transcription factor consisting of CLOCK (or NPAS2) and BMAL1. The gene products of per and cry accumulate in the cytoplasm, dimerize, enter the nucleus, and repress transactivation of the CLOCK-BMAL1 complex [14,15].

If this negative feedback loop were the entire story, one would anticipate only a single robust round of per/cry gene expression would occur, and then a steady, basal level of gene expression would persist. Rather, rhythmic relief of per/cry repression is achieved in part via progressive phosphorylation of PER by casein kinase 1 delta and epsilon, which targets the proteins for poly-ubiquitylation and degradation via the 26S proteasomal pathway [16,17].

Here, it should also be pointed out that this core molecular feedback loop is influenced by multiple clock-influenced feedback regulators, including kinases such as protein kinase A and the extracellular signal-related kinase (ERK)/mitogen-activated protein kinase (MAPK) cascade. Likewise, oscillations in transcriptional pathways mediated by the nuclear hormone receptor Rev-erb, the bZIP transcription factor E4BP4 [18] and the basic helix-loop-helix transcription factors Dec1 and Dec2 [19] modulate the amplitude and phasing of the clock and also serve as output pathways that impart circadian control over an array of biochemical and physiological processes [20,21].

Entrainment of the biological clock to external time cues appears to be mediated in part by rapid changes in the expression of per1 and per2. Along these lines, photic input drives a rapid increase in clock gene expression via an ERK/MAPK-cAMP response-element-binding protein (CREB) signaling cassette in the SCN [22] (Figure 1). Conversely, non-photic entrainment cues have been shown to trigger a rapid degradation of per1 and per2 expression [23]. As state variables of the clock, changes in the expression of per1 and per2 would lead to resetting of the molecular oscillator, and in turn re-entrainment of the circadian rhythm.

A deeper understanding of the functional significance of the circadian timing process was realized through gene-array-based analysis, which showed that large and diverse functional classes of genes within the SCN are under the control of the core-clock timing processes [24,25]. Likewise, robust rhythmicity in the periphery has been demonstrated for tissues, including the liver and heart [26-28]. A truly remarkable finding was the organ-specific nature of circadian gene expression profile. Hence, beyond the canonical core clock genes, only small subsets of genes have been found to oscillate in multiple organ systems; along these lines, Panda et al. [25] reported that 28 genes (out of approximately 650) were found to oscillate in both the SCN and the liver. These findings reveal that clock-controlled gene expression is tailored to match the physiological role of the tissue.

Circadian modulation of disease
The finding that circadian timing mechanisms are distributed through the body raises important questions about the relevance of the clock to human health and disease. Some of the most interesting work on this subject has utilized animal models to assess the physiological effects that disruption of timing has on disease progression. Along these lines, work by Filipski et al. [29] showed that destruction of the SCN accelerated malignant tumor growth by up to threefold. Likewise, experimental models that lead to a marked disruption of clock synchrony, such as constant light, and repetitive shifting of the light-dark cycle, have been shown to accelerate tumor growth and, relatedly, increase mortality rates [30-32]. In a telling parallel to these studies, a number of epidemiological studies have found that shift workers are at an elevated risk for several types of cancer, including breast, colorectal and prostate cancer [33-35]. Given work showing that the circadian clock modulates hormone output, immune cell function and cell division, the postulation that disruption of the circadian clock occurs as part of the disease process may not be surprising. However, many of these studies have also hinted at the possibility that disruption of the clock may be a key factor in the initiation of a disease state. In this regard, work in mouse models in which components of the core clock timing process are disrupted via genetic-based approaches have been particularly enlightening. Along these lines, Clock mutant mice are obese and hyperphagic, exhibiting hyperleptinemia, hepatic steatosis hyperglycemia and hypoinsulinemia [36]. In an interesting affirmation of the connection between the clock and metabolic disorders, SNP analysis of lean and obese humans has revealed an association between CLOCK polymorphisms and susceptibility to obesity [37]. In addition, haplotype analysis has identified CLOCK gene variants that may be protective against the development of obesity or that contribute to the development of metabolic syndrome, type 2 diabetes and cardiovascular disease [38]. It also appears that key components of the molecular clock are affected by alterations in metabolic activity. Along these lines, mice maintained on a high-fat diet, thus leading to obesity and metabolic syndromes, exhibited a marked alteration in core Clock gene expression within the liver and kidneys.
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http://genomemedicine.com/content/3/2/10

The SCN includes hormones, kinases, transcription factors and circadian clock. The clock-controlled transcriptome (CCT) within the transcriptome is under the direct or indirect influence of the cycling. In addition to the core clock genes, a large fraction of repression and, in turn, allowing for robust 24-hour clock gene transcription. Casein kinase 1 (CK1)-mediated phosphorylation production via suppression of CLOCK (CLK)- and BMAL1-mediated translocate to the nucleus and negatively regulate their own and cryptochrome (CRY) gene families. PERIOD and CRY dimers translocate to the nucleus and negatively regulate their own production via suppression of CLOCK (CLK)- and BMAL1-mediated transcription. Casein kinase 1 (CK1)-mediated phosphorylation triggers degradation of PER proteins, thus relieving transcriptional repression and, in turn, allowing for robust 24-hour clock gene cycling. In addition to the core clock genes, a large fraction of the transcriptome is under the direct or indirect influence of the circadian clock. The clock-controlled transcriptome (CCT) within the SCN includes hormones, kinases, transcription factors and microRNAs. Many of these gene products serve as phasing cues to ancillary oscillators, or function as feedback regulators of the core molecular clock.

Figure 1. Schematic overview of molecular circadian clock timing. (a) Suprachiasmatic nucleus (SCN; red circles) located within the ventral hypothalamus. The two nuclei serve as the master circadian clocks, orchestrating rhythmicity and phasing of ancillary clocks found in peripheral organ systems (c1) and throughout the brain (c2). Entrainment of the SCN clock to light is mediated by a direct projection from the retina. (b) The molecular clock feedback loop is centered on the rhythmic expression of the period (per1/2) and cryptochrome (cry1/2) gene families. PERIOD and CRY dimers translocate to the nucleus and negatively regulate their own production via suppression of CLOCK (CLK) and BMAL1-mediated transcription. Casein kinase 1 (CK1)-mediated phosphorylation triggers degradation of PER proteins, thus relieving transcriptional repression and, in turn, allowing for robust 24-hour clock gene cycling. In addition to the core clock genes, a large fraction of the transcriptome is under the direct or indirect influence of the circadian clock. The clock-controlled transcriptome (CCT) within the SCN includes hormones, kinases, transcription factors and microRNAs. Hence, one could envision an intertwining of genetic and lifestyle factors that impact metabolic fitness via alterations in circadian clock function.

Disruption of clock timing has also been shown to contribute to an array of nervous system disorders. Along these lines, familial advanced sleep phase syndrome (FASPS) is an autosomal dominant disorder characterized by a 3- to 4-hour advance in sleep onset and awakening [40]. Hence, an individual with FASPS may fall asleep at 7 pm rather than 11 pm, and with arousal at 3 am instead of 7 am. Linkage analysis revealed that FASPS was localized to a point mutation in the PER2 gene [41]. At a mechanistic level, this mutation is thought to eliminate a phosphorylation motif for casein kinase 1 epsilon, thus leading to excessive accumulations of PER2 and, in turn, an altered/advanced circadian cycle, initiating premature drive for sleep onset. Interestingly, dysregulation of the clock also appears to contribute to disturbances of the sleep-wake cycles in individuals with Alzheimer’s disease. These spontaneous night-time awakenings in patients with Alzheimer’s disease are one of the most challenging issues for caregivers [42]. Likewise, individuals suffering from Huntington’s disease exhibit clock-regulated disruption of neuroendocrine output, and of the sleep-wake cycles [43-45]. These examples are a mere shortlist of the profound functional roles that the clock plays in organismal physiology. The role of the clock in such diverse processes as cell division and xenobiotic responses, and its roles in regulation of behavior, cognition, arousal and aging are reviewed elsewhere [46-48].

Molecular control through microRNA

To further our understanding of the molecular underpinnings of the circadian clock, researchers have begun to explore the potential role of the non-coding class of small RNA species known as microRNAs. MicroRNAs are potent negative regulators of mRNA translation and, as such, play a central role in sculpting gene expression patterns. MicroRNAs are principally coded within introns and exons of non-coding RNA (that is, intergenic microRNA), and from introns of protein encoding genes (that is, introngenic microRNA) [49,50]. MicroRNAs are
frequently coded close to other microRNAs and transcribed as long RNA that contains multiple microRNAs in a cluster (polycistronic microRNAs) [50,51]. The biogenesis of microRNAs occurs via a multistep process, which is described in detail elsewhere [52-54]. In short, microRNAs are transcribed by RNA polymerase II as long single-stranded RNA, forming stem-loop (hairpin) structures, referred to as primary microRNAs. Primary microRNAs are processed to precursor microRNAs (55 to 70 nucleotides) by a protein complex of RNAase III enzyme Drosa and the double-stranded RNA-binding domain protein DGCR8/Pasha in the nucleus [55-57]. Precursor microRNAs are then exported to the cytoplasm where they are further processed to approximately 22 bp microRNA/microRNA* double-stranded species by a protein complex of RNAase III enzyme, Dicer, and double-stranded RNA-binding domain proteins TRBP/ PACT/loquacious [58,59]. Finally, microRNAs are loaded into the RNA-induced silencing complex, through which the gene silencing function of microRNAs occurs via a reduction in mRNA translation efficiency or mRNA stability [60-62].

**MicroRNA and clock timing**

Rhythmic control of microRNA expression appears to be conserved from plants to mammals, and has significant consequences for circadian timing. Along these lines, genome-wide tiling arrays have been used to profile the transcriptome of *Arabidopsis thaliana* as a function of the circadian cycle. Beyond the observation that approximately 25% of the coding genes exhibited circadian rhythms of expression, several rhythmically regulated microRNAs (that is, miR160B and miR167D) were identified [63]. This experiment was not designed to detect microRNA, and thus it is likely that the results underestimate the total number of oscillating microRNAs.

In addition to circadian control mechanisms, biologically relevant time cues, such as the daily light cycle, also drive rhythmic microRNA expression. Along these lines, *Arabidopsis* miR167, miR168, miR171 and miR398 have recently been shown to oscillate, with higher levels during the daytime than during the night [64] (Table 1). Interestingly, rhythmicity of these microRNAs was not governed by an internal circadian clock, suggesting photic control of some microRNAs. Within humans, it would not be surprising to observe a similar pattern of microRNA expression driven by the clock, exogenous time cues or both.

Circadian-regulated microRNA expression has also been described in *Drosophila melanogaster*. Array analysis of wild-type flies revealed that miR263a and miR263b showed robust circadian oscillations, which were absent in clock mutant flies [65] (Table 1). In a later study, Kadener *et al.* [66] found that abrogation of microRNA biogenesis led to both a marked rise in circadian-regulated gene expression and a disruption of circadian-regulated behavioral rhythms, thus indicating a role for microRNA in clock timing. AGO1 pull-down assays revealed that gene products involved in clock timing were targeted for microRNA-mediated degradation [64], indicating that core clock molecular feedback is not only regulated at a transcriptional and post-translational level, but also through post-transcriptional/microRNA-mediated control mechanisms.

In the mammalian SCN, microRNAs play a role in clock timing and entrainment. Cheng *et al.* [12] demonstrated that both miR132 and miR219 show oscillatory expression in wild-type mice, but not in circadian mutant mice. Both microRNAs have CRE-enhancer sequences allowing for CREB-dependent regulation [67], but only miR132 was shown to be inducible by light [12]. Importantly, miR219 appears to be under direct control of the molecular clock via the binding of CLOCK and BMAL1 to an E-box within its promoter region. In contrast, miR132 does not contain an E-box motif, and thus its rhythmicity is likely driven by an ancillary CREB-actuated transcriptional oscillator [12]. *In vivo* antisense silencing studies demonstrated that miR219 shortens the circadian period and that miR132 negatively regulates light-dependent resetting of the clock [12]. Both miR132 and miR219 affect *per1* expression, and thus influence the core circadian transcriptional loop [12,68]. However, neither miR132 nor miR219 appears to directly target *per1*; thus, the precise mechanisms by which these microRNAs affect the clock are not known.

**MicroRNA and peripheral circadian oscillators**

In addition to rhythmic microRNA expression in the SCN clock, peripheral oscillators also exert circadian control over microRNA expression. A microarray-based study of mouse liver revealed that over 13% of the probed microRNAs (85 out of 640) exhibited circadian expression patterns [69]. Target prediction analysis indicated that a number of cycling microRNAs were capable of binding to components of the core clock timing process and, for a subset of these microRNAs, there was an inverse expression pattern with their predicted clock gene targets. For example, expression of miR181d and its putative target CRY1 and its putative target BMAL1, were inversely correlated [69]. Conversely, miR328 and miR383 were positively correlated with the expression of their corresponding targets, *Per1* and *Cry1*. This unexpected relationship may be the result of a phase-lag in downregulation, or it may represent an example of positive microRNA regulation that has been uncovered in recent years [70].

In another study, a forward genetic screen was used to identify the miR192/194 gene cluster as inhibitors of the
per (per1 to 3) gene family [71]. Significantly, the miR192/194 cluster is highly expressed in the liver and kidneys (two oscillating organs) [72] and overexpression studies found that miR192/194 shortens the length of the circadian rhythm via a decrease in per gene expression.

The expression of miR122, a known regulator of liver metabolism [73], is also subject to circadian control. Although the mature strand microRNA does not oscillate, both primary and precursor miR122 show robust daily changes in expression, suggesting that transcription is under circadian control [74]. Knockdown of miR122 revealed a marked alteration in the expression of clock-controlled genes, many of which are involved in cholesterol and lipid metabolism. In an additional study, Kojima et al. [75] found that the rhythmic expression of the mRNA deadenylase Nocturnin is modulated by miR122 in the liver, suggesting that miR122 is a key intermediate in circadian regulation of hepatic metabolic activity.

Intestine physiology is also governed by the circadian oscillator, which synchronizes enterocyte proliferation and, in turn, nutrient absorbance. Within this context, it is interesting that miR16, miR20a and miR141 all oscillate in the intestine, and that miR16 regulates the rate of proliferation in jejunal crypts by suppressing essential G1/S cell cycle regulators [76]. Hence, microRNA may serve as a link in the circadian control of intestinal physiology.

Finally, several microRNAs have been found to exhibit diurnal oscillations in the mouse retina, including miR96, miR124a, miR103, miR182, miR106b, miR422a and miR422b [77]. Of these, both miR96 and miR182 show potential targeting and downregulation of the clock-regulated gene adenylyl cyclase 6. In a separate study, miR26a, which is regulated by both CLOCK and CREB, was found to drive circadian rhythms in the production of the L-type voltage-gated calcium channel α1C subunit in cone photoreceptors [78]. This observation is of functional significance, given the central role that L-type calcium channels play in rhythmic melatonin synthesis and, in turn, circadian retinal physiology [79,80]. Finally, in humans, a polymorphism in pre-miR182 (which exhibits diurnal rhythmicity in the retina [77]) is significantly associated with major depression [81]. Relative to wild-type miR182, mutant miR182 was found to trigger a marked reduction in the expression of the circadian gene Clock. These data raise the possibility that mutant miR182 contributes to circadian clock ailments associated with major depression [82].

**Chronotherapeutic approaches and future outlook**

The finding that microRNA affects key clock timing processes (that is, the core molecular oscillator, clock entrainment and clock output) raises the prospect that chronotherapeutic approaches designed to affect microRNA physiology might prove efficacious.

Along these lines, some of the most tractable ailments associated with clock physiology may result from a damped SCN oscillator, which is observed as part of the aging process [83-85]. Hence, approaches to increase the robustness of the clock timing process may offset fragmentation of the sleep-wake cycle, which is often observed as a function of aging. This approach could also prove beneficial to individuals suffering from age-regulated neurodegenerative disorders, such as Alzheimer’s disease. Therapeutic approaches could take the form of microRNA antisense or ‘sponge’ constructs [86], designed to target particular microRNAs, and in turn limit their ability to suppress components of the core clock timing process, or clock output genes. Of note, in mouse models, microRNA knockdown using cholesterol-conjugated 2- O-methyl RNA antisense (that is, antagonirs [87]) has already been used to assess the functional roles of miR132 and miR219 in SCN clock physiology [12]. Conversely, transgenic RNA interference-based approaches designed to harness the endogenous microRNA processing machinery could be used to selectively uncouple an aberrant clock-timing gene or clock-output gene from a specific physiological process. This approach could prove particularly useful as part of a cancer treatment strategy.

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**Table 1. Light and clock-controlled microRNAs**

| Model          | Organ                      | MicroRNAs                        | Reference(s) |
|----------------|----------------------------|----------------------------------|--------------|
| Cell culture   |                            | miR328, miR383                   | [70]         |
| Arabidopsis    |                            | miR160, miR167, miR168, miR171, miR398 | [63,64]      |
| Drosophila     |                            | miR263                           | [65]         |
| Chicken        | Photoreceptors             | miR26a                           | [78]         |
| Rat            | Intestine                  | miR16, miR20a, miR141            | [76]         |
| Mouse          | Liver                      | miR122, miR181, miR191           | [69,73-75]   |
|                | Retina                     | miR96, miR103, miR106, miR124, miR182, miR422 | [77]         |
|                | Suprachiasmatic nucleus    | miR219, miR132                   | [12]         |
| Human          |                            | miR182                           | [81]         |
Obviously, there are numerous challenges that will need to be addressed for these types of approaches to be implemented. For example, the inability of RNA constructs (for example, antagonism and sponges) to cross the blood-brain barrier is a significant impediment to minimally invasive therapeutic strategies. Further, issues related to the timing of antisense administration and organ-specific therapeutic targeting will need to be addressed. At the least, the practical application of these methods will require a deeper understanding of microRNA biology, the clock, and its role in the regulation of a myriad of gene networks that contribute to human physiology and pathology.

**Abbreviations**
- CREB, cAMP response-element-binding protein
- ERK, extracellular signal-related kinase
- FASPS, familial advanced sleep phase syndrome
- MAPK, mitogen-activated protein kinase
- SCN, suprachiasmatic nucleus
- SNP, single-nucleotide polymorphism

**Competing interests**
The authors declare that they have no competing interests.

**Authors' contributions**
All three authors contributed to the writing of the manuscript.

**Acknowledgements**
This work was supported by grants MH62335, NS066345 and NS067409 from the National Institutes of Health.

**Published:** 17 February 2011

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doi:10.1186/gm224

Cite this article as: Hansen KE et al. MicroRNAs: a potential interface between the circadian clock and human health. *Genome Medicine* 2011, 3:10.