Circadian Rhythms in Environmental Health Sciences

Jacqueline M. Leung 1 · Micaela E. Martinez 1

© The Author(s) 2020

Abstract
Purpose of Review This review aims to explore how circadian rhythms influence disease susceptibility and potentially modify the effect of environmental exposures. We aimed to identify biomarkers commonly used in environmental health research that have also been the subject of chronobiology studies, in order to review circadian rhythms of relevance to environmental health and determine if time-of-day is an important factor to consider in environmental health studies. Moreover, we discuss opportunities for studying how environmental exposures may interact with circadian rhythms to structure disease pathology and etiology.

Recent Findings In recent years, the study of circadian rhythms in mammals has flourished. Animal models revealed that all body tissues have circadian rhythms. In humans, circadian rhythms were also shown to exist at multiple levels of organization: molecular, cellular, and physiological processes, including responding to oxidative stress, cell trafficking, and sex hormone production, respectively. Together, these rhythms are an essential component of human physiology and can shape an individual’s susceptibility and response to disease.

Summary Circadian rhythms are relatively unexplored in environmental health research. However, circadian clocks control many physiological and behavioral processes that impact exposure pathways and disease systems. We believe this review will motivate new studies of (i) the impact of exposures on circadian rhythms, (ii) how circadian rhythms modify the effect of environmental exposures, and (iii) how time-of-day impacts our ability to observe the body’s response to exposure.

Keywords Circadian rhythms · Environmental health · Biomarkers · DNA methylation · Asthma · Breast cancer

Introduction
Circadian rhythms exist in species throughout the tree of life, from single-cell organisms (e.g., cyanobacteria and Trypanosomes) to humans. Circadian rhythms drive 24-h cycles in physiology and behavior and evolved in response to predictable changes in Earth’s environment that occur around the day-night cycle [1]. By structuring biological processes into 24-h cycles, the circadian system enables organisms to synchronize their internal biology with their external environment and optimally time activities to maximize fitness and survival [2]. For example, circadian rhythms in human metabolic processes are believed to be evolutionarily advantageous for structuring metabolic activity according to the human diel lifestyle of nighttime sleep, daytime wake activity, and daytime food intake [3].

In mammals, including humans, circadian rhythms are present in nearly all tissues and cells in the body [4]. The master circadian clock, in the suprachiasmatic nucleus (SCN) of the hypothalamus, is composed of clock genes. Clock genes include several positive genes (i.e., CLOCK, BMAL1) and negative genes (i.e., PER, CRY) that are organized through cell-autonomous transcription-translation feedback loops [5]. The CLOCK:BMAL1 heterodimer binds to DNA and drives the rhythmic transcription of PER and CRY, whose protein products in turn inhibit the activity of the CLOCK:BMAL1 complex. Together, these factors regulate the downstream rhythmic expression of thousands of clock-controlled genes that generate oscillations in physiology and behavior. Light entrains the SCN and keeps clock genes
appropriately transcriptionally aligned with the day-night cycle [6]. The SCN then helps to entrain peripheral clocks throughout the body via neuronal signals, hormones, metabolites, and changes in body temperature [7].

The maintenance of circadian rhythms is critical to human health and survival. There are deleterious perturbations to circadian rhythms that include circadian disruption, physical and social jetlag (social jetlag occurs due to different sleep/wake schedules kept on work versus non-work days), and circadian misalignment. Circadian disruption occurs due to a lack of synchronization between body clocks and the day-night cycle and can cause misalignment among the various circadian clocks present throughout the body [8]. Well-known causes of circadian disruption/misalignment include various occupational and household environmental factors such as shift work [9], jetlag [10], and exposure to artificial light-at-night [11]. Overall, circadian disruption, whether due to jetlag or shift work, is mainly driven by light exposure during the body’s biological night (i.e., subjective night) [12]. There is a growing literature demonstrating that chronic circadian disruption can contribute to the development of diseases, including asthma, cancer, metabolic syndromes, and cardiovascular disease [3, 13, 14]. Disease risk associated with circadian disruption led the American Medical Association in 2012 to adopt a policy statement concluding that “The natural 24-hour cycle of light and dark helps maintain precise alignment of circadian biological rhythms...Pervasive use of nighttime lighting disrupts these endogenous processes and creates potentially harmful health effects and/or hazardous situations with varying degrees of harm” [12, 15], and, in 2007, the World Health Organization’s International Agency for Research on Cancer (IARC) classified shift work that involves circadian disruption as a probable human carcinogen [16].

There is a growing literature demonstrating that many diseases, including asthma and heart attack, have important circadian features, sometimes with the incidence and/or severity varying across the 24-h period [1, 8, 13, 17]. For example, the frequency of sudden cardiac death peaks in the morning [18], with a 40% increased risk of heart attack between 6 a.m. and noon [17], and the severity of asthma symptoms often worsens at night [19]. A better understanding of how the circadian system influences disease—particularly how circadian rhythms modify the body’s response to the environment and exposures—could provide new insights into disease etiology and lead to new strategies for prevention and treatment. Here, we aim to identify chronobiology studies that may inform environmental health research relating to exposures such as endocrine-disrupting chemicals, air pollution, noise/light pollution, and environmental determinants of chronic disease. We discuss circadian impacts on the pathophysiology and/or treatment of asthma, heart disease, and breast cancer because these are diseases for which both chronobiology and environmental health research exist to date, and we explore how circadian rhythms influence biological targets of environmental exposures and biomarkers that are commonly used in the study of environmental exposures (Fig. 1). As a future perspective, we also discuss ways in which studies of environmental exposures and environmental determinants of health can deploy circadian and time-dependent sampling to study how circadian rhythms may act as a modifier of disease.

Circadian Rhythms and Disease

In reviewing the chronobiology literature, we identified three diseases—asthma, heart disease, and breast cancer—that...
have been the subject of chronobiology studies and have also been studied in environmental health in relation to environmental exposures and/or environmental determinants of health. The intersection of environmental health and chronobiology likely go beyond these diseases, but this is the state of the field to date.

The incidence and severity of many diseases varies across the day and night [1, 8]. We describe how circadian rhythms in underlying biological processes can shape the overall presentation of disease (e.g., as in asthma and heart disease) and how circadian disruption due to occupational environmental exposures may elevate risk of disease (e.g., as in breast cancer).

**Asthma**

One of the major diseases known to have circadian variation is asthma. Asthma is a chronic, inflammatory disease of the respiratory system associated with exposure to environmental factors such as allergens, smoke, and air pollution [20]. Importantly, asthma symptoms show pronounced time-of-day variation in occurrence and severity. For instance, exacerbated asthma symptoms occur primarily at night and peak around 04:00 a.m. [19]. The majority of asthma deaths also occur between midnight and 06:00 a.m. [21]. The elevated nighttime incidence of asthma exacerbations has been associated with circadian changes in pulmonary function, including a nocturnal increase in airway inflammation, bronchial responsiveness, and eosinophils (i.e., cells of the immune system associated with asthma and allergy) in bronchial tissue [22]. The nocturnal increase in eosinophils is correlated with an increase in lung airway obstruction. Animal models of asthma have indicated that the expression of the clock gene BMAL1 plays a role in eosinophil trafficking and the production of the cytokine IL-5 [23], which is a target of multiple asthma drugs and has been shown to be under epigenetic regulation [24].

Not only do experimental and clinical studies suggest the pathology of asthma is impacted by circadian rhythms, these rhythms are likely important for disease manifestation and treatment. For instance, individuals with asthma often have circadian rhythms with an exaggerated amplitude. Fluctuations in airway obstruction, as measured by peak expiratory flow (PEF) over 24 h, were found to be synchronized between healthy and asthmatic individuals, with both having more obstruction occurring at night. However, the amplitude of PEF variation was 50.9% greater in asthma patients compared to controls [25]. From a pathology perspective, we hypothesize that the amplitude of an individual’s circadian rhythm in PEF could in part predispose them to asthma attacks and/or modify severity when an attack is triggered, for instance by environmental exposures. Figure 2 illustrates how circadian amplitude and time-of-day (i.e., circadian phase) could physiologically position individuals such that they are susceptible to an asthma exacerbation. Since circadian rhythms are present in utero, with humans having circadian rhythms as early as 30 weeks of gestation [26], they have the potential to play a role in diseases such as asthma that may stem from developmental periods.

![Fig. 2 Hypothetical mechanism by which circadian rhythms in inflammation affect asthmatic symptoms. The figure shows a theoretical threshold of inflammation in bronchial tissue, above which an individual may clinically present with asthma. First, we hypothesize high amplitude circadian rhythms (e.g., individual 1) may predispose individuals to asthma. Second, we hypothesize that time-of-day effects on asthma are due to circadian rhythms opening up a window of susceptibility that modifies the effect of environmental exposures, resulting in a higher likelihood of asthma exacerbations](image-url)
Heart Disease

Exposure to environmental pollutants, such as pesticides, tobacco smoke, and fine particulate matter, play an important role in the development and severity of cardiovascular disease [27, 28]. These exposures impact aspects of physiology that are known to affect cardiovascular risk, including changes in blood pressure, endothelial function, blood lipids, and blood coagulation markers [29]. Importantly, these physiological aspects also display time-dependent oscillations throughout the day-night cycle [30]. For instance, blood pressure surges upon waking [31], and vascular endothelial function is impaired in the morning [32]. Circadian variation in these factors parallel the pattern of adverse cardiovascular events, including myocardial infarction [33, 34], stroke [35], ventricular arrhythmia [36], and sudden cardiac deaths [18, 37], with increased numbers typically observed in the morning between 06:00 a.m. and 12:00 p.m. [38].

Disruptions of the molecular clock have also been shown to contribute to cardiovascular disease. Knockout of the BMAL1 gene causes dilated cardiomyopathy in mouse models [39], and cardiac-specific deletion of BMAL1 initiates diastolic dysfunction, increases fibrotic responses, and impairs resolution of inflammation [40], thereby reducing survival from cardiomyopathy [41]. BMAL1 gene deletion in cardiomyocytes also impacts cardiac ion homeostasis that is important in the electrical activity of the heart by abating circadian expression of Na+ and K+ channels, which could lead to sudden cardiac death [42, 43]. Together, these data indicate that circadian rhythms play an important role in structuring cardiovascular disease risk. The mechanism by which circadian rhythms generate time-of-day susceptibility to acute cardiovascular events remains to be revealed. However, due to circadian rhythms in the biomarkers used to assess cardiovascular disease (i.e., blood pressure, lipid panels, etc.), it is important to control for time-of-day when evaluating such clinical biomarkers. Additionally, since some exposures may vary in presence and intensity throughout the day-night cycle (e.g., air pollution), the interplay between cycles in exposure and circadian rhythms should also be considered.

Breast Cancer

The disease for which circadian rhythms have been most intensely studied in environmental health is breast cancer. In 2006, using data from The Nurses’ Health Study II, which included over 100,000 female nurses from the U.S., it was discovered that women who reported 20+ years of night shift work had a relative risk of breast cancer of 1.79 compared to women who did not report shift work [44]. It has been hypothesized that circadian disruption caused by shift work is responsible for elevated risk. There is now further evidence implicating circadian disruption in breast cancer risk; specifically, women in the Nurses’ Health Study II living in the most highly light polluted areas in the U.S. had a 14% higher risk of breast cancer compared to women living in the least light polluted areas [45]. A recent case-control study in Spain also found that both breast cancer and prostate cancer risk were associated with artificial light-at-night in the blue light spectrum, with blue being the most important part of the spectrum for circadian rhythms [46]. It is hypothesized that light pollution in cities contributes to light-at-night exposure on the individual level; therefore, light pollution can be used as a proxy for disruptive light-at-night exposure. The mechanisms by which light-at-night and circadian disruption may elevate cancer risk remain unknown; however, it may be related to the disruption of rhythms in the neuroendocrine and/or immune system that, when intact, may confer protection.

There are many limitations to current epidemiological studies of circadian rhythms and cancer. Light-at-night exposure and circadian disruption are difficult to measure, and there are potentially many confounding variables associated with shift work, light-at-night, and cancer. One recent study of over 100,000 women in the United Kingdom did not find consistent associations between breast cancer risk and self-reported night shift work in the past 10 years [47]. Taken together, the studies of cancer and circadian disruption highlight the need to develop protocols for evaluating circadian rhythm disruption as a risk modifier for diseases with environmental influence, such as breast cancer. These studies also highlight the need for including light pollution in the portfolio of environmental exposures that may impact health.

Circadian Control of Clinical and Biological Biomarkers in Environmental Health

Environmental exposures have the ability to modify biological processes and mechanisms that contribute to downstream health and disease risks. At the same time, many of these biomarkers vary throughout the day-night cycle and have the potential to make individuals more or less vulnerable to environmental insults depending on the time of exposure. Here, we briefly discuss various processes known to have circadian rhythms and explore the mechanisms that link these processes to environmental exposures and health. We focus on five types of biological processes that are often used as markers of environmental exposures: oxidative stress, inflammation, endocrine disruption, DNA methylation, and histone modifications, all of which are important in epigenetic research and studies of exposure-related disease risk [48].

Oxidative Stress

Various environmental pollutants are able to generate free radical reactions that result in oxidative damage. Environmental
exposures, such as UV light or chemical pollutants, can lead to the production of reactive oxygen species (ROS), which can result in tissue injury, DNA damage, and cell death [49]. Protective enzymes and small-molecule antioxidants can counteract the detrimental effects of ROS that would otherwise leave cells in a prolonged state of oxidative stress [50].

Oxidative stress is often measured by assessing DNA damage, lipid peroxidation, and protein oxidation, all three of which have been shown to display circadian rhythms. It has been suggested that rhythms in oxidative stress biomarkers are due to rhythms in the response to oxidative stress, as opposed to rhythms in exposure, because these oscillations parallel the daily activity rhythms of protective enzymes and antioxidants [51]. Rodent models have been used in the study of oxidative stress rhythms. In rats, superoxide dismutases, which catalyze the dismutation of O$_2$− into the less reactive species O$_2$ and H$_2$O$_2$, have been shown to peak at night in the cerebral cortex [52]. Additionally, daily oscillations of glutathione (GSH), an antioxidant that removes hydroperoxides and neutralizes ROS, peaks in the middle of the light phase in rats and is inversely related to lipid peroxidation, suggesting that a decrease in GSH activity during the night could be partly responsible for the peak in lipid peroxidation at night [52]. Various other antioxidants and protective enzymes have also been shown to undergo daily redox cycles [53, 54].

In addition to the characterization of oxidative stress rhythms in humans and rodent models, studies in circadian clock mutants of Drosophila have demonstrated circadian control of antioxidant expression. In Drosophila, ROS levels, protein carbonylation, and mortality were significantly higher in wild-type flies exposed to H$_2$O$_2$-induced oxidative stress during the day versus the night, and this daily susceptibility rhythm was abolished in flies with a null mutation in the clock gene PER [55]. Additionally, mice deficient in the clock gene BMAL1 had an upregulation of ROS levels in several tissues. There is evidence that the circadian clock is important for ROS homeostasis and the aging process because the upregulation of ROS was associated with accelerated aging, and the administration of the antioxidant N-acetyl-L-cysteine ameliorated the premature aging [56, 57]. Taken together, studies of circadian rhythms in oxidative stress responses reveal that environmental health studies of oxidative stress should not only account for time-of-day when evaluating biological biomarkers, but could also evaluate circadian rhythms of participants in longer term studies of oxidative stress, DNA damage, and aging.

**Inflammation**

Over the last decade, studies of circadian rhythms in the mammalian immune system have flourished. There are very well documented, self-sustained rhythms in the immune system driven by clock genes expressed within cells of the immune system. Because these rhythms are controlled by gene expression outside of the master clock in the SCN, the clock of the immune system is considered to be a peripheral clock. One of the earliest discoveries in this area was that leukocytes display circadian trafficking throughout the body. For instance, in mice—which are nocturnal and whose rhythms are expected to be phase shifted from diurnal mammals like humans—leukocytes migrate from the blood and enter into tissues at night. This trafficking has been shown to involve signals from the autonomic nervous system [58] and can directly impact the outcome of inflammatory diseases [59]. Particularly, circadian control of the immune system can dictate the timing of inflammatory disease onset [60, 61]. For example, rheumatoid arthritis patients exhibit increased joint inflammation, pain, and stiffness in the early morning hours, which results from nighttime synthesis and release of proinflammatory cytokines and chemokines, leading to increased cell migration to inflamed tissues during this time [62].

Circadian rhythms in the immune system and inflammatory response influence the intensity and onset of inflammation triggered in response to environmental stimuli. Exposure to environmental contaminants, such as allergens and air pollution, has the ability to cause inflammation; therefore, underlying rhythms in inflammation have the potential to modify the effect of inflammatory exposures. It is therefore necessary to understand the circadian regulation of inflammatory pathways in order to understand the pathophysiology of inflammatory diseases and appropriately time treatment.

**Endocrine Disruption**

Endocrine-disrupting chemicals (EDCs), such as pesticides, flame retardants, plastics, and pharmaceuticals, are mostly man-made chemicals released into the environment that can interfere with the body’s hormonal signaling pathways and cause adverse health effects [63]. EDCs can alter the normal functioning of the endocrine system by mimicking natural hormones in the body, blocking the interaction of hormones with their receptors, or interfering with the way hormones or their receptors are made [64]. Certain classes of EDCs, including phthalates, dichlorodiphenyltrichloroethane (DDT), and bisphenol A (BPA), can affect reproductive health by mimicking or blocking the effects of male and female sex hormones [65]. For instance, interquartile range increases in the phthalate di-2-ethylhexyl phthalate (DEHP) detected in urine are associated with a 7.84% decrease in serum testosterone in young men and a 24.0–34.1% decrease in boys [66]. Due to the long recognized circadian rhythm in testosterone, models in such studies adjust for time-of-day when measuring the EDC effect on testosterone. In addition to testosterone, many hormones have been shown to have circadian rhythms in production.
In men, plasma testosterone peaks in the morning around 08:00 a.m. and decreases to its lowest concentrations in the evening [67, 68]. Total, free, and bioavailable testosterones are on average 30–35% higher at 08:00 a.m. compared to mid and late afternoon levels in young men [69]. The amplitude of the circadian rhythm declines with age, dropping to only a 10% difference between morning and late day in men 70 years of age [69]. The strong circadian rhythm in testosterone highlights the importance of accounting for rhythms when evaluating the effect of an environmental exposure, or for evaluating a biomarker in individuals of different ages. For example, early studies investigating the relationship between serum testosterone levels and age found inconsistent results [70-73], likely due to differences in blood sampling time; however, when controlling for sampling time, an age-related decrease in testosterone levels was found in blood taken in the morning in young versus older men, but not in samples taken in the afternoon, which resulted from the loss of circadian rhythmicity in testosterone with aging [68]. With a 30–35% higher plasma testosterone level in the morning verses late day in young men, and a 7.84–34.1% decrease in urinary testosterone in boys and young men exposed to DEHP, it is clear that the natural daily variation in testosterone is comparable to the level of variation due to the environmental exposure, thus requiring the time-of-day adjustment in models.

DNA Methylation

The study of DNA methylation is currently a major area of Environmental Health Sciences. Environmental pollutants have been linked to disease phenotypes through epigenetic modifications such as DNA methylation, histone modifications, and microRNAs [77]. DNA methylation occurs post-DNA replication and involves the addition of a methyl group at the fifth carbon position of cytosine residues, which leads to changes in gene expression without changing the underlying DNA sequence. Aberrant DNA methylation has been observed in various cancer types [78], autoimmune diseases [79], metabolic disorders [80], and aging [81]. Methylation analysis therefore has the potential to aid in the diagnosis and treatment of various diseases [82].

In contrast to the prevailing view that DNA methylation is a relatively stable epigenetic marker, recent evidence suggests that DNA methylation can fluctuate daily and seasonally [83, 84]. Levels of DNA methylation in a number of gene promoter regions have been shown to display circadian changes [85, 86]. In mice, expression of enzymes involved in DNA methylation and demethylation in the SCN can be induced by light in a circadian phase-dependent manner, which could be a way that the SCN drives circadian clock plasticity [83]. Daily cycles in DNA methylation have also been demonstrated in the blood of healthy individuals, with increased levels of DNA methylation occurring at night [87]. Work in human cells lines has revealed that DNA methylation sites can fluctuate within a single cell cycle [85, 86, 88], thereby demonstrating the dynamic nature of DNA methylation. Although one may assume daily cycles in DNA methylation could be driven by daily cycles in cell proliferation and associated DNA replication, slow proliferating cells of the liver, lung, and brain in mice also exhibit daily cycles in methylation, particularly at CpG sites [89-91]. These circadian oscillations of cytosine modifications are also prevalent in human neutrophils and are associated with complex diseases including leukemia, schizophrenia, and diabetes [92].

Histone Post-translational Modifications

Histones are nuclear globular proteins that undergo posttranslational modifications such as acetylation, methylation, phosphorylation, SUMOylation, ribosylation, and ubiquitination [93]. Histones alter chromatin structure and hence modulate gene expression and genome stability [94, 95]. Environmental pollutants, such as pesticides and metals, can affect histone modifications and/or alter the function of enzymes involved in histone modifications [96, 97]. For example, arsenic exposure can inhibit histone deacetylases and induce chromatin opening by histone hyperacetylation [98]. Aberrant patterns of histone posttranslational modifications in the genome have been associated with various human diseases, including autoimmune...
diseases [99], neurodegenerative diseases [100], and cancer [101], and thus represent an important epigenetic alteration in environmental diseases.

A direct link between the circadian clock and epigenetics via histone modifications can be found in the CLOCK gene, which has been shown to possess intrinsic histone acetyltransferase activity that is necessary for circadian gene expression [102]. CLOCK also acetylates non-histone substrates, such as BMAL1, in a circadian manner [103], suggesting that other targets in the body may also be affected. Indeed, CLOCK can also acetylate the glucocorticoid receptor, thereby attenuating its DNA binding and regulating its function [104, 105]. The activation of several clock-controlled genes, including PER1, PER2, and Dbp, has also been associated with rhythmic changes in histone acetylation at their promoters [106–108]. These studies suggest that one way in which the circadian clock exerts transcriptional control is through mechanisms involving histone modification and chromatin remodeling. Daily alterations in these factors have been linked to circadian changes in processes such as metabolism and cellular proliferation [109, 110]. The study of circadian histone modification has largely focused on the expression of clock genes; importantly, however, mouse models have also shown that 43% of all protein coding genes have circadian expression somewhere in the body (e.g., liver, kidney, lung, brown fat, etc.) [111•]; therefore, circadian histone modification could be impacting the expression of genes in addition to the core clock genes.

**Conclusion**

In conclusion, circadian rhythms are a main structuring feature of the human body and exist at multiple levels of organization from molecular to physiological processes. Circadian rhythms impact molecular processes within cells, such as responses to oxidative stress, DNA replication, DNA methylation, histone modification, and gene expression. As for cellular and physiological processes, circadian rhythms drive cell trafficking in the immune system and the production of neuroendocrine and reproductive hormones. We suggest that all studies should aim to control for circadian rhythms when selecting biomarkers of exposure or health. Biological processes and biomarkers vary in the degree to which they display rhythmicity, and those that are rhythmic vary in their amplitude of circadian variation. When measuring the effect of exposures on biological processes, it is important to know the magnitude of circadian variation relative to the amount of variation due to environmental exposures. Lack of controls for circadian rhythms could mask important variation due to exposures.

When considering exposures such as air pollution, endocrine-disrupting chemicals, etc., it is important to recognize that the time-of-day of exposure could impact health risk. For example, exposure to artificial light at night can impact biological processes and have pathological consequences, whereas artificial light exposure during the day would not have the same effect [12•]. Given that the circadian clock is entrained by light, studies should acknowledge that the rhythmicity of certain biomarkers may be affected when sampled under varying lighting conditions, such as in the winter (short days) vs. summer (long days) or for people who spend most of their time indoors (under artificial light) vs. outdoors (under natural light). Ideally, but often unfeasible, study designs should deploy circadian and time-dependent sampling, with multiple samples being collected at regular intervals across several days. Such longitudinal sampling allows researchers to better assess within-individual changes over time and can indicate whether we are missing important variation in physiological and behavioral responses. For instance, this can be accomplished through the collection of at least three sampling timepoints within a 24-h period: shortly after waking, midday, and immediately before bed. If this is not feasible and a study is limited to the collection of a single sample, researchers should control for the time in which the sample is collected or include that information in the subsequent statistical analyses.

In this review, we have highlighted the importance of circadian rhythms in environmentally influenced diseases such as asthma, breast cancer, and heart disease. We believe that the intersection of circadian biology and environmental health offers a novel angle for exploring the mechanism of disease pathology and susceptibility. In particular, one interesting avenue for future research is determining how circadian rhythms may influence individual susceptibility to environmental exposures throughout the day. For instance, would the same dose of PM$_{2.5}$ air pollution have the same effect on the body in the morning as the evening? If there is circadian variation in susceptibility, should this be taken into account in air pollution policy? Studies could also evaluate how light-at-night, which causes circadian rhythm disruption, interacts with other environmental exposures such as noise and air pollution, potentially magnifying health effects. Lastly, the majority of best-selling drugs and essential medicines target the product of circadian genes [111•]. This alone demonstrates the importance of acknowledging rhythms in the overall science of health, as we seek to improve treatment and prevention.

**Funding Information** Drs. Micaela Martinez and Jacqueline Leung are funded by the NIH Directors Early Independence Award. Research reported in this publication was supported by the Office of the Director, National Institutes of Health, under Award Number DP5OD023100. This publication was also supported by a pilot grant from the NIEHS Center for Environmental Health in Northern Manhattan ES009089.

**Compliance with Ethical Standards**

**Conflict of Interest** Dr. Martinez and Dr. Leung declare no conflict of interest.
Human and Animal Rights and Informed Consent  This article does not contain any studies with human or animal subjects performed by any of the authors.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

References

Papers of particular interest, published recently, have been highlighted as:

- Of importance

1. Martinez-Bakker M, Helm B. The influence of biological rhythms on host-parasite interactions. Trends Ecol Evol. 2015;30:314–26.
2. Yerushalmi S, Green RM. Evidence for the adaptive significance of circadian rhythms. Ecol Lett. 2009;12:970–81.
3. Davies SK, Ang JE, Revell VL, Holmes B, Mann A, Robertson FP, et al. Effect of sleep deprivation on the human metabolome. Proc Natl Acad Sci U S A. 2014;111:10761–6.
4. Dibner C, Schibler U, Albrecht U. The mammalian circadian timing system: organization and coordination of central and peripheral clocks. Annu Rev Physiol. 2010;72:517–49. This is a good review discussing the structure and function of the mammalian circadian timing system.
5. Takahashi JS. Transcriptional architecture of the mammalian circadian clock. Nat Rev Genet Nature Publishing Group. 2017;18:164–79.
6. Reppert SM, Weaver DR. Coordination of circadian timing in mammals. Nature. 2002;418:935–41.
7. Albrecht U. Timing to perfection: the biology of central and peripheral circadian clocks. Neuron. 2012;74:246–60.
8. Roenneberg T, Merrow M. The circadian clock and human health. Curr Biol. 2016;26:R432–43.
9. Rajaratnam SMW, Howard ME, Grunstein RR. Sleep loss and circadian disruption in shift work: health burden and management. Med J Aust. 2013;199:S11–5.
10. Sack RL. Jet Lag. N Engl J Med. 2010;362:440–7.
11. Lunn RM, Blaske DE, Coogan AN, Figueiro MG, Gorman MR, Hall JE, et al. Health consequences of electric lighting practices in the modern world: a report on the National Toxicology Program’s workshop on shift work at night, artificial light at night, and circadian disruption. Sci Total Environ. Elsevier B.V. 2017;607:590–608;1073–84.
12. Stevens RG, Brainard GC, Blaske DE, Lockey SW, Motta ME. Adverse health effects of nighttime lighting: comments on an American medical association policy statement. Am J Prev Med. 2013;45:343–6. This article summarizes a major policy statement that was adopted by the American Medical Association in 2012 regarding the potential harmful effects of nighttime lighting on human health.
13. Martin RJ, Banks-Schlegel S. Chronobiology of asthma. Am J Respir Crit Care Med. 1998;158:1002–7.
14. Evans JA, Davidson AJ. Health consequences of circadian disruption in humans and animal models. Prog Mol Biol Transl Sci. 2013.
15. Blask D, Brainard G, Gibbons R, Lockey S, Motta M. Report 4 of the Council on Science and Public Health (A-12). Light pollution: adverse health effects of nighttime lighting. Rep 4 Counc Sci Public Heal. 2012;4–5.
16. Stevens RG, Hansen J, Costa G, Haus E, Kauppinen T, Aronson KJ, et al. Considerations of circadian impact for defining “shift work” in cancer studies: IARC working group report. Occup Environ Med. 2011;68:154–62.
17. Elliott W. Cyclic and circadian variations in cardiovascular events. Am J Hypertens. 2001;14:S291–5.
18. Muller JE, Ludmer PL, Willich SN, Tofler GH, Aylmer G, Klangos I, et al. Circadian variation in the frequency of sudden cardiac death. Circulation. 1987;75:131–8.
19. Bohadana AB, Hannhart B, Teculescu DB. Nocturnal worsening of asthma and sleep-disordered breathing. J Asthma. 2002;39:85–100.
20. Gilmour MI, Jaakkola MS, London SJ, Nel AE, Rogers CA. How exposure to environmental tobacco smoke, outdoor air pollutants, and increased pollen burdens influences the incidence of asthma. Environ Health Perspect. 2006;114:627–33.
21. Hetzel MR, Clark TJH, Branthwaite MA. Asthma: analysis of sudden deaths and ventilatory arrests in hospital. Br Med J. 1977;1:808–11.
22. Kraft M, Martin RJ, Wilson S, Djkunavic R, Holgate ST. Lymphocyte and eosinophil influx into alveolar tissue in nocturnal asthma. Am J Respir Crit Care Med. 1999;159:228–34.
23. Zaslona Z, Case S, Early JO, Lalor SJ, McLaughlin RM, Curtis AM, et al. The circadian protein BMAL1 in myeloid cells is a negative regulator of allergic asthma. Am J Physiol Lung Cell Mol Physiol. 2017;312:L855–60.
24. Peng C, Cardenas A, Rifas-Shiman SL, Hivert MF, Gold DR, Platts-Mills TA, et al. Epigenome-wide association study of total serum immunoglobulin E in children: a life course approach. Clin Epigenetics. 2018;10:1–14.
25. Hetzel MR, Clark TJH. Comparison of normal and asthmatic circadian rhythms in peak expiratory flow rate. Thorax. 1980;35:732–8.
26. Mirrman M, Kok JH, Boer K, Wolf H. Perinatal development of human circadian rhythms: role of the foetal biological clock. Neurosci Biobehav Rev. 1992;16:371–8.
27. Brook RD, Rajagopalan S, Pope CA, Brook JR, Bhatnagar A, Diez-Roux AV, et al. Particulate matter air pollution and cardiovascular disease: an update to the scientific statement from the american heart association. Circulation. 2010;121:2331–78.
28. Csordas A, Bernhard D. The biology behind the atherothrombotic effects of cigarette smoke. Nat Rev Cardiol. Nature Publishing Group. 2013;10:219–30.
29. Costello KE, Navas-Acien A, Kaufman JD. Environmental factors in cardiovascular disease. Nat Rev Cardiol. Nature Publishing Group. 2015;12:627–42.
30. Thosar SS, Butler MP, Shea SA. Role of the circadian system in cardiovascular disease. J Clin Invest. 2018;128:2157–67.
31. Millar-Craig MW, Bishop CN, Raftery EB. Considerations of circadian impact for defining “shift work” in cancer studies: IARC working group report. Occup Environ Med. 2011;68:154–62.
32. Otto ME, Svatikova A, De Mattos Barretto RB, Santos S, Hoffmann M, Khandheria B, et al. Early morning attenuation of endothelial function in healthy humans. Circulation. 2004;109:2507–10.
33. Goldberg RJ, Brady P, Muller JE, Chen Z, de Groot M, Zonneveld P, et al. Time of onset of symptoms of acute myocardial infarction. Am J Cardiol. 1990;66:140–4.
34. Muller JE, Stone PH, Turi ZG, Rutherford JD, Czeisler CA, Parker C, et al. Circadian variation in the frequency of onset of acute myocardial infarction. N Engl J Med. 1985;313:1315–22.
35. Elliott WJ. Circadian variation in the timing of stroke onset: a meta-analysis. Stroke. 1998;29:992–6.
36. Twidale N, Taylor S, Heddle WF, Ayres BF, Tonkin AM. Morning increase in the time of onset of sustained ventricular tachycardia. Am J Cardiol. 1989;64:1204–6.
37. Willich SN, Goldberg RJ, Maclure M, Perriello L, Muller JE. Increased onset of sudden cardiac death in the first three hours after awakening. Am J Cardiol. 1992;70:65–8.
38. Muller JE, Toller GH, Stone PH. Circadian variation and triggers of onset of acute cardiovascular disease. Circulation. 1989;79:733–43.
39. Lefla M, Campbell KS, Fung HY, Jin JP, Esser KA. Development of dilated cardiomyopathy in Bmal1-deficient mice. Am J Physiol Heart Circ Physiol. 2012;303:475–85.
40. Ingle KA, Kain V, Goel M, Prabhu SD, Young ME, Halade GV. Cardiomyocyte-specific Bmal1 deletion in mice triggers diastolic dysfunction, extracellular matrix response, and impaired regulation of inflammation. Am J Physiol Heart Circ Physiol. 2015;309:H1827–36.
41. Young ME, Brewer RA, Pellicari-Garcia RA, Collins HE, He L, Birky TL, et al. Cardiomyocyte-specific BMAL1 plays critical roles in metabolism, signaling, and maintenance of contractile function of the heart. J Biol Rhythm. 2014;29:257–76.
42. Schroder EA, Lefla M, Zhang X, Bartos D, Fung HY, Zhao Y, et al. The cardiomyocyte molecular clock, regulation of Scn5a, and arrhythmia susceptibility. Am J Physiol Cell Physiol. 2013;304:954–65.
43. Schroder EA, Burgess DE, Zhang X, Lefla M, Smith JL, Patwardhan A, et al. The cardiomyocyte molecular clock regulates the circadian expression of Kcnh2 and contributes to ventricular repolarization. Heart Rhythm. Elsevier. 2015;12:1306–14.
44. Schernhammer ES, Kroenke CH, Laden F, Hankinson SE. Night work and risk of breast cancer. Epidemiology. 2006;17:108–11.
45. This study shows a link between rotating night shift work and breast cancer risk and is the first prospective report on this association in premenopausal women.
46. James P, Bertrand KA, Hart JE, Schernhammer ES, Tamimi RM, Laden F. Outdoor light at night and breast cancer incidence in the nurses’ health study II. Environ Health Perspect. 2017;125:087010.
47. Garcia-Saenz A, de Miguel AS, Espinosa M, Atzori LM, Amoros M, Llorca J, et al. Evaluating the association between artificial light-at-night exposure and breast and prostate cancer incidence in the generations study cohort. Br J Cancer. 2019;121:172–81.
48. Schernhammer ES, Kroenke CH, Laden F, Hankinson SE. Night work and risk of breast cancer. Epidemiology. 2006;17:108–11.
49. Schernhammer ES, Kroenke CH, Laden F, Hankinson SE. Night work and risk of breast cancer. Epidemiology. 2006;17:108–11.
50. Matés JM, Birky TL, et al. Cardiomyocyte-specific BMAL1 plays critical roles in metabolism, signaling, and maintenance of contractile function of the heart. J Biol Rhythm. 2014;29:257–76.
51. Silver AC, Arjona A, Walker WE, Wikir E. The circadian clock controls toll-like receptor 9-mediated innate and adaptive immunity. Immunology. 2012;36:251–61.
52. Cutolo M, Straub RH, Buttigerei F. Circadian rhythms of nocturnal hormones in rheumatoid arthritis: translation from bench to bedside. Ann Rheum Dis. 2008;67:905–8.
53. Druzd D, Matveeva O, Ince L, Harrison U, We H, Schmal C, et al. Adrenergic nerves govern cardiovascular leukocyte recruitment to tissues. Immunity. 2012;36:251–61.
54. Schernhammer ES, Kroenke CH, Laden F, Hankinson SE. Night work and risk of breast cancer. Epidemiology. 2006;17:108–11.
55. Schegar TT, Johnson AF, Birmbaum LS, Colborn T, Guillette LJ, Crews DP, et al. Minireview: endocrine disruptors: past lessons and future directions. Mol Endocrinol. 2016;30:833–47.
56. Schegar TT, Janesick A, Blumberg B, Heindel JJ. Endocrine disrupting chemicals and disease susceptibility. J Steroid Biochem Mol Biol. 2011;127:204–15.
57. Meeker JD, Ferguson K, Underdahl J. Delayed puberty and the risk of breast cancer: a systematic review of the literature. Clin Endocrinol. 2016;85:528–37.
58. Reinberg A, Lagoquy M. Circadian and circannual rhythms in sexual activity and plasma hormones (FSH, LH, testosterone) of five human males. Arch Sex Behav. 1978;7:13–30.
59. Brenner WJ, Vitiello MV, Prinz PN. Loss of circadian rhythmicity in blood testosterone levels with aging in normal men. J Clin Endocrinol Metab. 1983;56:1278–81.
60. Brambilla DJ, Matsumoto AR, Araujo AB. Menopause and risk of breast cancer. J Clin Endocrinol Metab. 2014;99:4346–52.
61. Reinberg A, Lagoquy M. Circadian and circannual rhythms in sexual activity and plasma hormones (FSH, LH, testosterone) of five human males. Arch Sex Behav. 1978;7:13–30.
62. Brenner WJ, Vitiello MV, Prinz PN. Loss of circadian rhythmicity in blood testosterone levels with aging in normal men. J Clin Endocrinol Metab. 1983;56:1278–81.
63. Brambilla DJ, Matsumoto AR, Araujo AB. Menopause and risk of breast cancer. J Clin Endocrinol Metab. 2014;99:4346–52.
73. Harman SM, Tsitouras PD. Reproductive hormones in aging men. I. Measurement of sex steroids, basal luteinizing hormone, and Leydig cell response to human chorionic gonadotropin. J Clin Endocrinol Metab. 1980;51:35–40.

74. Rahman SA, Grant LK, Gooley JW, Rajaratnam SMW, Cezisler CA, Lockley SW. Endogenous circadian regulation of female reproductive hormones. J Clin Endocrinol Metab. 2019;104:6049–59.

75. Mortola JF, Laughlin GA, Yen SS. A circadian rhythm of serum follicle-stimulating hormone in women. J Clin Endocrinol Metab. 1992;75:816–4.

76. Pollack AZ, Schisterman EF, Goldman LR, Mumford SL, Albert CA, Lockley SW. Variations in DNA methylation patterns during the cell cycle of HeLa cells. Epigenetics. 2007;2:54–65.

77. Brown SE, Fraga MF, Weaver IC, Birdasco M, Szfy M. Variations in DNA methylation patterns during the cell cycle of HeLa cells. Epigenetics. 2007;2:54–65.

78. Kangaspeska S, Stride B, Metivier R, Polycarpou-Schwarz M, Ilberson D, Carmouche RP, et al. Transient cyclical methylation of promoter DNA. Nature. 2008;452:112–5.

79. Metivier R, Gallais R, Tiffonc C, Loron P, Zurekowska RZ, Carmouche RP, et al. Cyclical DNA methylation of a transcriptionally active promoter. Nature. 2008;452:45–50.

80. Bönsch D, Hothorn T, Krieglstein C, Koch M, Nehmer C, Lenz B, et al. Daily variations of homocysteine concentration may influence circadian function via changes in global transcription and DNA methylation. This epigenetic modification may thus provide a mechanism for light-dependent circadian clock function.

81. Stevenson TJ, Prendergast BJ. Reversible DNA methylation regulates seasonal photoperiodic time measurement. Proc Natl Acad Sci U S A. 2013;110:16615–19.

82. Kansgaspesa S, Stride B, Metivier R, Polycarpou-Schwarz M, Ilberson D, Carmouche RP, et al. Transient cyclical methylation of promoter DNA. Nature. 2008;452:112–5.

83. Metivier R, Gallais R, Tifféno C, Le Périon C, Zurkowski RZ, Carmouche RP, et al. Cyclic DNA methylation of a transcriptionally active promoter. Nature. 2008;452:45–50.

84. Bönsch D, Hothorn T, Krieglstein C, Koch M, Nehmer C, Lenz B, et al. Daily variations of homocysteine concentration may influence methylation of DNA in normal healthy individuals. Chronobiol Int. 2007;24:315–25.

85. Brown SE, Fraga MF, Weaver IC, Birdasco M, Szfy M. Variations in DNA methylation patterns during the cell cycle of HeLa cells. Epigenetics. 2007;2:54–65.

86. Oh G, Ebrahimi S, Carlucci M, Zhang A, Nair A, Groot DE, et al. Cytosine modifications exhibit circadian oscillations that are involved in epigenetic diversity and aging. Nat Commun. 2018;9:1–11.

87. Coulson RL, Yasui DH, Dunaway KW, Laufer BI, Vogel C, Ciernia B, et al. Daily variations of homocysteine concentration may influence circadian function via changes in global transcription and DNA methylation. This epigenetic modification may thus provide a mechanism for light-dependent circadian clock function.

88. Pollack AZ, Schisterman EF, Goldman LR, Mumford SL, Albert CA, Lockley SW. Variations in DNA methylation patterns during the cell cycle of HeLa cells. Epigenetics. 2007;2:54–65.

89. Oh G, Ebrahimi S, Carlucci M, Zhang A, Nair A, Groot DE, et al. Cytosine modifications exhibit circadian oscillations that are involved in epigenetic diversity and aging. Nat Commun. 2018;9:1–11.

90. Lim ASP, Klein HU, Yu L, Chibnik LB, Ali S, Xu X, et al. Diurnal and seasonal molecular rhythms in human neocortex and their relation to Alzheimer’s disease. Nat Commun. Nature Publishing Group. 2017;8.

91. Oh G, Konecnirovics K, Ebrahimi S, Carlucci M, Groot DE, Nair A, et al. Circadian oscillations of cytosine modification in humans contribute to epigenetic variability, aging, and complex disease state.

Publisher's Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.