Chapter

Consequences of Artificial Light at Night: The Linkage between Chasing Darkness Away and Epigenetic Modifications

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Abstract

Epigenetics is an important tool for understanding the relation between environmental exposures and cellular functions, including metabolic and proliferative responses. At our research center, we have devolved a mouse model for characterizing the relation between exposure to artificial light at night (ALAN) and both global DNA methylation (GDM) and breast cancer. Generally, the model describes a close association between ALAN and cancer responses. Cancer responses are eminent at all light spectra, with the prevalent manifestation at the shorter end of the visible spectrum. ALAN-induced pineal melatonin suppression is the principal candidate mechanism mediating the environmental exposure at the molecular level by eliciting aberrant GDM modifications. The carcinogenic potential of ALAN can be ameliorated in mice by exogenous melatonin treatment. In contrast to BALB/c mice, humans are diurnal species, and thus, it is of great interest to evaluate the ALAN-melatonin-GDM nexus also in a diurnal mouse model. The fat sand rat (Psammomys obesus) provides an appropriate model as its responses to photoperiod are comparable to humans. Interestingly, melatonin and thyroxin have opposite effects on GDM levels in P. obesus. Melatonin, GDM levels, and even thyroxin may be utilized as novel biomarkers for detection, staging, therapy, and prevention of breast cancer progression.

Keywords: melatonin, thyroxin, light-at-night, global DNA methylation, diurnal species, breast cancer, biomarkers

1. Introduction

Since the invention of electrical light in 1879 by Thomas Alva Edison, artificial light at night (ALAN) has become a definitive feature of human development with accelerated increase concurrent with urbanization and industrialization. The light emitted from the original bulb of Edison known as incandescent bulb was weak, with a dominant long wavelength emission above 560 nm. Most of the incandescent electrical energy is dissipated as heat energy, thus making this type of illumination energetically inefficient. Therefore, new illumination technologies were developed, in order to discover efficient bulbs that transfer most of the electrical energy into light. White fluorescent and light-emitting diodes (LED) are examples of energy efficient bulbs developed to decrease carbon dioxide production from electric power plants, thus lessening the greenhouse effect. One of the adverse outcomes of using efficient
illumination at night time is the emission of shorter wavelengths (SWLs) that further exacerbate the health and ecological problems associated with a new source of environmental pollution currently known as ALAN [1–3]. Light pollution is increasing rapidly, resulting in a more illuminated world, where outdoor and indoor illumination sources are increasing ALAN in developed and developing countries [4, 5].

From an anthropological perspective, electric light has brought pronounced benefits including advancing urbanization and industrialization by increasing productivity, but we are also increasingly being aware of serious public health and ecological negative impacts emerging from disrupting the adaptive temporal organization of biological responses [6–8]. Certainly, multiple studies have shown the effects of light pollution on social, behavioral, physiological, and molecular responses in many different taxa, including insects [9], fishes [10], amphibians [11], reptiles [12], birds [13], and mammals [14], as well as plants [15]. Some of the most disturbing effects of ALAN on health are metabolic dysfunction and cancer progression [2, 16]. In mice and humans, several lines of evidence suggest a close association between ALAN levels and both obesity and breast cancer progression [17–19]. Here, we focus on ALAN as a novel environmental polluter that disrupts biological timing (temporal organization) and consequently may provoke severe health risk, particularly breast cancer development through epigenetic modifications. First, the mammalian photoperiodic system is reviewed in relation to light perception and downstream endocrine responses for timing biological rhythms. Thereafter, we discuss the sensitivity of the photoperiodic system to the spectral composition of ALAN, particularly SWL illuminations. We further discuss the ALAN signal transduction pathway involved in melatonin suppression and aberrant epigenetic modifications in breast cancer progression. Therefore, melatonin and epigenetics are suggested as new biomarkers for breast cancer prevention. Finally, melatonin and thyroxin treatments in the diurnal fat sand rat (Psammomys obesus) are discussed in relation to their potential role in mediating the environmental exposures at the molecular level via epigenetic modifications, particularly global DNA methylation (GDM).

2. The mammalian photoperiodic system

In an early study, it has been demonstrated that the blind mole rat (Spalax ehrenbergi) responded differently to short and long photoperiod manipulations in regard to its capability to cope with low ambient temperature exposure [20]. Results of a more recent study on S. ehrenbergi manifested robust and differential responses in metabolism, stress, and melatonin levels to ALAN of different spectral compositions and acclimation duration [21]. These results suggested that the vestigial retina of this species still expresses photoreceptors that are involved mainly in nonvisual response. Currently, the mammalian eye is described as a dual-function organ, expressing photoreceptors for both visual and nonvisual responses [22]. The visual response is mediated by two distinct photoreceptor types, rods and cones, which control scotopic vision and photopic vision, respectively [23]. The nonvisual responses are mainly mediated by intrinsically photosensitive retinal ganglion cells (ipRGCs) that express the photopigment melanopsin. Even though the ipRGCs are connected with rods and cones by bipolar cells, they mediate nonvisual responses including photo-entrainment of biological rhythms [24].

First, photoperiodic signals are perceived by ipRGCs that express the photopigment melanopsin [25]. The detected environmental light signal by the ipRGCs synchronizes the master circadian clock located in the mammalian hypothalamic suprachiasmatic nucleus (SCN) by the retinohypothalamic tract (RHT). The
SCN regulates the synthesis and release of the hormone melatonin by the pineal gland through multiunit sympathetic nerves from the superior cervical ganglion (SCG). The SCG presynaptic sympathetic terminals release noradrenalin that interacts with postsynaptic α- and β-adrenergic receptors to regulate synthesis and release of pineal melatonin [26]. In mammals, the activity of the adrenergic SCG terminals that innervate the pineal gland is stimulated by darkness and inhibited by light [27]. Under dark conditions, stimulation of the pineal adrenergic receptors increases cellular cAMP levels leading to the activation of aryl-alkyl-amine-N-acetyltransferase (AA-NAT), a key enzyme in melatonin synthesis [28]. The nocturnal increase in the enzymatic activity of AA-NAT is strongly inhibited by light exposure, consequently leading to a rapid decrease in nocturnal melatonin levels [29]. The pinealocytes are the primary neuroendocrine cells that synthesis melatonin by sequential hydroxylation and decarboxylation of its precursor tryptophan to serotonin. Thereafter, serotonin is acetylated by the rate-limiting enzyme AA-NAT and methylated by the enzyme hydroxyindole-O-methyltransferase (HIOMT) to the final product of melatonin [28, 30]. Finally, the activity of both AA-NAT and HIOMT is under photoperiodic control at the transcriptional level showing distinct diurnal rhythms with peak levels during night and nadir levels during the day [31].

3. Melatonin suppression as an indicator of SWL pollution

In most mammals, no level of light exposure is powerless regarding melatonin suppression and even low intensity and short-term exposures can reduce its production and lead to decreased circulating levels [32, 33]. Nonetheless, melatonin suppression is strongly wavelength- and irradiance-dependent, with faster and more robust response at the SWL end of the visible spectrum below 500 nm [19, 34, 35]. A large-scale study comparing the effect of different light technologies on melatonin production in humans demonstrated that the strongest suppression occurred in response to 4000 and 5000 K LED lights compared with incandescent, halogen, and fluorescent counterpart lightening systems [36]. Narrow bandwidth blue LED exposure (λ = 469 nm, ½ peak bandwidth = 26 nm) decreased melatonin levels in an irradiance dose-dependent manner, and this light was more effective in decreasing the hormone levels compared with that of 4000 K of white fluorescent at twice the energy of the latter [37]. In horses, 1 h exposure of 3 lux SWL blue light (468 nm) administered only to one eye was sufficient to decrease melatonin levels compared with control animals [38]. Furthermore, blue LED pulses (2-s pulse every 1 min for 1 h, λ = 450 nm) administrated through closed human eyelids markedly suppressed nocturnal melatonin levels and delayed the melatonin onset phase [39–41]. While the eyelids can weaken irradiance and wavelength ([42], light signals can still penetrate them, be detected by the retinal photoreceptors, and affect circadian regulation [43]. In humans, blue LED exposure (40 lux, 470 nm) emitted from display screens (tablets and computers), suppressed nocturnal melatonin in a duration-dependent manner [44, 45] and melatonin suppression showed higher sensitivity to wavelength compared with intensity manipulations [46].

Together, it is clear that the adverse effects of light pollution are strongly manifested by the SWL portion of the spectrum. As the LED illumination is becoming ubiquitous in every aspect of our modern life, the expected increase in light pollution may exacerbate the problem since higher irradiance and shorter wavelengths would be emitted by the energy efficient technology [47, 48]. Accordingly, the American Medical Association [49] passed a resolution in 2016 calling upon
Communities in the USA to avoid using LED lighting in public domains as it is enriched with SWL [49]. In summary, SWL-ALAN is a source of pollution and should be removed from public spaces through legislation.

4. ALAN as an environmental change and a model for studying epigenetic modifications

The flexibility and the sensitivity of the endocrine system play an adaptive role in determining the success and survival of organisms under contentiously changing environmental conditions in their habitat [50]. As the endocrine system regulates several functions, it is expected to be the first system to respond to environmental changes such as ALAN by coordinating body functions to maintain homeostasis during the exposure. The core stimulus-response of the endocrine system to ALAN relies on four main components, including the pineal gland, the hypothalamic-pituitary-gonadal (HPG) axis, the hypothalamic-pituitary-thyroid axis (HPT), and the hypothalamic-pituitary-adrenal (HPA) axis [51]. The elaborated hormonal responses generated by these axes to ALAN exposure might be mediated by transcriptional regulation of gene expression via epigenetic modifications [52]. Therefore, epigenetic-elicited alteration in gene expression is a potential transduction pathway by which hormonal responses (e.g., melatonin) may mediate environmental exposures (e.g., ALAN). Conversely, the ALAN-induced alteration in melatonin rhythms may also exert endocrine responses via epigenetic modifications [53].

The incidences of breast and prostate cancers show close association with light pollution particularly in urbanized and industrialized regions [2, 54]. Several epidemiological studies have found direct association between light pollution and incidence of breast cancer in women as well as prostate cancer in men [18, 55, 56]. Furthermore, the strong association between light pollution and cancer incidences displays divergent spatial disruption with higher incidences in urban compared with rural regions [57, 58]. Evidence for direct association between ALAN and cancer development comes also from animal studies.

In rats, ALAN exposure accelerated the growth rates of induced-tumors, including mammary cancer [59–62]. Studies under control conditions demonstrated that 30-min ALAN per midnight emitted from either white fluorescent or blue LED illuminations can accelerate tumor growth and lung metastatic activity in female BALB/c mice inoculated with 4T1 mammary carcinoma [63, 64]. Indeed, the effects of ALAN on tumor growth have been demonstrated at different spectral compositions with markedly higher cancer burden in response to lighting exposure lower than 500 nm [19].

These studies have related the increased cancer burden to aberrant epigenetic modifications, particularly advanced global DNA hypo-methylation. Promoter hyper-methylation of cancer suppressor genes and global DNA hypo-methylation are characterizing epigenetic patterns in breast cancer cells [65, 66]. These aberrant epigenetic modifications may contribute to increase cancer burden by eliciting genomic instability and activation of both oncogenes and metastatic related genes, as well as silencing tumor suppressor genes. Generally, prominent decreased methylation in repetitive DNA elements is a common trait in most cancer cells [67]. Demethylation of pro-metastatic genes is normally suppressed by DNA methylation and might advance gene overexpression leading to genetic instability that increases the risk of developing cancer [68, 69]. DNA hypomethylation can be detected at an early stage of breast cancer and is correlated with the degree of tumor differentiation [70, 71]. Altogether, the close association between aberrant
DAN hypomethylation and tumorigenesis, particularly of breast cancer, is well-established, but the underlying mechanism remains poorly understood, especially how the adverse ALAN effects are mediated.

5. Melatonin as a mediating signal linking ALAN and epigenetic-induced cancer

Since the melatonin hypothesis was first proposed during the late twentieth century by Stevens [72], multiple studies in human and nonhuman animals have provided direct and indirect evidence that melatonin suppression by ALAN could impose health risks, including metabolic disorders and cancer progression [2, 54]. The importance of melatonin in the regulation of several biological functions depends heavily on its lipophilic and hydrophilic traits that make it omnipresent in all cell compartments, principally in the nucleus [73]. Indeed, low levels of 6-sulfatoxymelatonin (6-SMT), the major metabolites of the hormone in urine [74], have been demonstrated to correlate with increased risk of breast cancer in postmenopausal women [75–77]. Furthermore, women with blindness or long sleep duration (elevated melatonin levels) present reduced breast cancer risk relative to normal women [78, 79].

Physiological blood concentration of melatonin blocked human leiomyosarcoma (soft tissue sarcoma) proliferation by inhibiting tumor metabolic and genetic pathways presumable by suppression of cellular cAMP levels via melatonin receptor [80]. In hepatocellular carcinoma-induced mice, melatonin treatment suppressed tumor cell proliferation through arresting the cell cycle [81]. The metastatic activity of oral squamous cell carcinoma was notably reduced by melatonin-mediated inhibition of tumor-associated neutrophils [82], inflammatory cells involved in promoting several solid tumors [83]. Similarly, the anti-oncogenic property of melatonin has been demonstrated also in other cancer types, including lung [84], gastric [85], ovarian [86], and colon [87], as well as breast cancers [88].

Melatonin could mediate its effects of cancer development via epigenetic modifications, particularly GDM [89]. Melatonin treatment to MCF-7 cell lines significantly increased DNA methylation that was associated with increased transcriptional levels of the tumor metastasis suppressor gene glypican-3 and decreased expression levels of the oncogenes EGR3 and POU4F2/Brn-3b [90]. In estrogen-receptor-related breast cancer, melatonin may decrease transcriptional levels of the aromatase gene (involved in the regulation of estrogen synthesis) by either methylation of the gene or deacetylation of the promoter gene [91]. Additionally, nocturnal melatonin treatment can rectify the induced DNA demethylation, tumor growth, and metastatic activity by both blue LED and fluorescent ALAN in 4T1 mammary cancer cell-inoculated female BALB/c mice [63, 64]. In a more recent study that evaluated the effects of ALAN and melatonin treatment at different spectral compositions in 4T1-inoculated BALB/c mice, a tissue-specific response in GDM was detected [19]. In this study, the tumor tissue manifested the most prominent changes in GDM showing an inverse wavelength-dependent correlation that was reversed by melatonin. Conversely, other tissues (e.g., lung, liver, and spleen) showed mixed results of positive, negative, or indifferent correlation between methylation levels and both wavelength and melatonin treatments [19]. Largely, melatonin may regulate epigenetic modifications in a number of tumor-related genes mainly by DNA methylation, but other modifications are also possible.

The strong association between ALAN, DNA hypo-methylation, and melatonin suppression may be of significant clinical importance. DNA methylation and melatonin can be utilized as biomarkers for detecting and preventing breast cancer development. The traditional diagnosis method for breast cancer is scanning by
mammography, which is a useful technique to identify the growth of cancer. The mammography cannot predict risk for breast cancer as it indicates its existence, but trends in melatonin suppression and DNA methylation can provide a simple, noninvasive, and reliable tool for predicting cancer risk, particularly among a group of high-risk individuals for developing the disease such as night shift workers. Bearing in mind that epigenetic modifications are reversible [92], early treatment by melatonin or any other analogs [93] for individuals at high risk can be very effective in preventing breast cancer. We are aware today, that genetics factors such as breast cancer genes are not the major causes of the malignancy and other external factors are heavily involved. Therefore, much more attention should be given to environmental changes that link endocrinology with epigenetic modifications.

Collectively, in diurnal humans, circadian disruption enforced by activity impinging on the inactive period during the nighttime is recurrently associated with a number of health problems. However, a direct link between ALAN-induced circadian disruption and health risks is still difficult to clearly establish as most data are derived from epidemiological and nocturnal animal studies [94]. Therefore, integrating diurnal animal models of chronodisruption with epidemiological and nocturnal model studies would add a significant value in defining potential direct signal transduction pathways mediating the environmental exposure impacts on physiology and health. Consequently, we conducted a preliminary study to investigate the effects of hormonal manipulations in diurnal species on physiological and epigenetic regulations. This preliminary study is a first step in a large-scale study using diurnal mouse model to elucidate the association between ALAN-induced circadian disruption and the development of health problems at the behavioral, physiological, and molecular levels.

6. Physiological and epigenetic responses to melatonin and thyroxin in diurnal species

Bearing in mind that humans are diurnal, understanding the physiological and epigenetic response to ALAN in human disease can benefit significantly from using a diurnal species such as the fat sand rat (*Psammomys obesus*). This species is a good model because it is a photoperiodic species that responds to photoperiod with robust daily rhythms in a number of physiological functions, including body temperature, melatonin levels, and AA-NAT activity [95, 96]. Furthermore, *P. obesus* is a useful model for studying human health and diseases such as metabolic disorders, obesity, diabetes, inflammation, and cardiovascular impairment [97–100]. Since most previous studies on photoperiodic responses were conducted on nocturnal species, in our research center at the University of Haifa, we use *P. obesus* as a model for studying photoperiodic and hormonal manipulations. In *P. obesus*, melatonin and body temperature rhythms were diminished in response to constant dim blue light exposure, while melatonin treatment restored the disrupted rhythms [63]. Although the previous studies have clearly indicated that as a diurnal species, *P. obesus* can respond to photoperiod and light manipulations, the underlying mechanism mediating the effect of the environmental changes remains unknown. An unanswered question is how melatonin and thyroxin interact to mediated environmental-induced epigenetic modifications. To answer this question, male *P. obesus* were acclimated to a long photoperiod cycle of 16L:8D at an ambient temperature of 24 ± 1°C and humidity of 45 ± 2%. Lights during the day were emitted from cool fluorescent lamps at 470 lux and 470 nm. Rats were caged individually and provided with ad libitum tap water and low energy diet. At the end of 3-week acclimation period, rats were either untreated, *i.p.* injected
with melatonin, thyroxin, or melatonin and thyroxin in combination 3 h after the dark period onset (01:00 h). Hormones were daily administered for 3 weeks at a dose of 50 μg/kg for melatonin and 2 mg/kg for thyroxin. During the experimental period, body mass ($W_b$) was monitored every other day and urine samples were collected by a noninvasive method [19] at 4 h intervals over a 28 h period. Urine samples were used to measure the major metabolite of melatonin in urine, 6-SMT [101]. The urinary metabolite concentrations were assayed by enzyme-linked immunosorbent assay utilizing a commercial IBL kit (RE54031) following the manufacturer's protocol. Finally, digit tips were collected from rats at the end of urine collection for DNA isolation (High pure PCR Template Preparation Kit, Roche) and subsequently for GDM analysis (MethylFlash™ Methylated DNA Quantification Kit, Epigentek). All experimental procedures were performed with the approval from the Ethics and Animal Care Committee of the University of Haifa.

The results showed that melatonin alone significantly increased $W_b$ from day 1 compared with controls, but with a decreasing magnitude with time (Figure 1A). Mass gain on day 1 was approximately 1.5-fold higher compared with that at the last. T4 also increased $W_b$ from day 1 to day 5 compared with controls, but with significantly lesser effect compared with melatonin. Thereafter, mass was decreased showing a moderate mass loss from day 13 to day 21 compared with controls. Thyroxin and melatonin in combination markedly decreased $W_b$ with time compared with all other groups. Mass gain decreased from 0.46 ± 0.88% at day 1 to −20.21 ± 2.56% at day 21. Thyroxin can regulate $W_b$ by increasing heat production through nonshivering thermogenesis by changing membrane permeability to sodium, increasing the pump activity to maintain cell homeostasis in brown adipose tissue, resulting in higher body temperature values and loss in $W_b$ [102].

Melatonin may operate through increasing the amount of brown adipose tissue, thus increasing heat production by increasing energy expenditure. Melatonin and thyroxin in combination provoked considerably more mass loss than melatonin alone, suggesting that melatonin may act synergistically with thyroxin to evoke mass loss in rats, due to the combined effect of increasing energy expenditure.

Body temperature rhythms were notably altered only in response to T4 treatment, while melatonin alone and in combination with thyroxin had no effect on body temperature compared with controls (Figure 1B). Furthermore, the significant decrease in body temperature following treatments with thyroxin and melatonin in combination, compared with T4 alone, suggests that melatonin and thyroxin exert a significant antagonistic effect on body temperature.

Thyroxin treatment had no effect on mean 6-SMT levels but altered the daily rhythms with higher amplitude and delayed acrophase by approximately 2 h (Figure 2A). Finally, melatonin treatment elicited hypomethylation while thyroxin alone or thyroxin and melatonin in combination exerted comparable effects on GDM levels showing marked hypermethylation compared with control levels (Figure 2B). Similar to $W_b$, thyroxin and melatonin may have exerted synergistic effects on promoting DNA hypermethylation, but this effect did not reach statistical significance.

These results suggest that melatonin and thyroxin have a role in the regulation of body temperature and apparently metabolism, in which the former may attenuate metabolism and the latter may accelerate it. Both hormones exerted inverse effects on global DNA levels, suggesting that different transduction pathways are involved in the circadian regulation of body temperature in P. obesus. The results suggest also that change in body temperature is more sensitive to thyroxin treatment than melatonin, as the effect of the latter was masked in the combined treatment with the other hormone.
However, in humans, melatonin may interact with the HPT axis to modulate the circadian rhythm of body temperature [104]. In mammals, the HPT axis plays a major role in several adaptive functions such as growth, development, metabolic rate, thermogenesis, heart rate, immune, and reproductive responses [105]. The HPT releasing and stimulating hormones as well as the thyroid hormones (T4 and T3) are under photoperiodic control presumably by the pars tuberalis of the adenohypophysis [106, 107]. In rats, T3 and T4 concentrations exhibit significant circadian rhythms with elevated levels during the dark period compared with the counterpart light period [108]. The nocturnal increase in the thyroid hormones was reported also in the rat pineal gland following an increase in type I 5′-iodothyronine deiodinase activity, which catalyzes the conversion of T4 to T3 [109]. Furthermore, the thyroid hormones are crucial photoperiodic regulators of several physiological

Figure 1.
Percentage change in body mass (A) and body temperature (B) in long-day acclimated P. obesus under four conditions: control no treatments, thyroxin (T4) treatment, melatonin (MLT) treatment, and combined treatment with T4 + MLT. Data are presented as mean ± standard error of nine animals. Different letters represent statistically significant difference among groups (Bonferroni, P < 0.01). # vs. day 21 (Bonferroni, P < 0.02).
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processes including energy metabolism and reproduction [110, 111]. While the relation between the HPT axis and the photoperiodic system are well-characterized, there are limited studies on the effect of ALAN on the HPT axis. However, due to the link with the photoperiodic system, environmental perturbation of the circadian clock by ALAN is expected to alter the activity of the HPT axis, including the thyroid hormones. In hamsters under short-day photoperiod, low levels of ALAN elevated the levels of thyroid-stimulating-hormone (TSH) receptors causing advanced Wb and gonadal growth [112]. Continuous exposure to ALAN decreased

Figure 2.
Daily rhythms of urinary 6-sulfatoxymelatonin (A) and global DNA methylation (B) levels in long-day acclimated P. obesus under four conditions: control no treatments, thyroxin (T4) treatment, melatonin (MLT) treatment, and combined treatment with T4 + MLT. In panel A, the best-fitted cosine curve (black and gray lines) and Cosinor estimates (period, P-value, and percentage of the rhythm [PR]) are depicted [103]. The gray area in each plot represents the length of the dark period. Data are presented as mean ± standard error of seven to nine animals. Different letters represent statistically significant difference among groups (Bonferroni, P < 0.01).
TSH, but increased both T3 and T4 in mice [113]. In birds, long-term exposure to ALAN increased both the blood levels of the thyroid hormones and W [114]. Overall, ALAN may induce aberrant epigenetic modifications by disrupting endocrine axes such as HPT axis that interacts with melatonin to manifest the adverse effects of the environmental exposure. However, the exact mechanism of action by which HPT axis may directly, or via melatonin, mediate the disruption effects of ALAN on the circadian system and promote downstream health risk is still unclear, and further efforts are warranted for elucidating it.

7. Conclusions

Currently, it is clear that electric light not only has remarkable anthropological advantages, but also severe adverse ecological and public health concerns. One of the most alerting impacts of ALAN on public health is the potential association between SWL exposure and cancer development, particularly in urbanized regions worldwide. ALAN effects are suggested to be mediated at the cellular level by inducing epigenetic modifications via nocturnal melatonin suppression. A schematic of ALAN-induced adverse effects is presented in Figure 3. Accordingly, light signals including ALAN are detected by ipRGCs and conveyed to the SCN by RHT. During a normal light dark cycle, melatonin is synthesized and secreted to the blood during the night, where it entrains central and peripheral oscillators to regulate normal physiological responses. Conversely, ALAN suppresses melatonin levels causing chronodisruption and misalignment in central and peripheral oscillators resulting in impaired physiological responses. The central and peripheral oscillators can be regulated directly by the melatonin signal or indirectly by modifying the body temperature rhythms [115]. In mice, daily variations in body temperature rhythms have been demonstrated to synchronize circadian gene expressions [116] and these central-controlled variations can be utilized to regulate variant peripheral circadian clocks in mammals [117]. Consequently, in diurnal species, thyroxin as an endocrine pathway is presumably involved in center circadian regulation of peripheral clocks by modifying body temperature daily rhythms.

These effects are presumably mediated by aberrant epigenetic modifications. Therefore, DNA methylations, which are a reversible modification in genes, triggered by melatonin, are a promising mechanism linking between environmental exposures like ALAN and hormonal/cellular pathway mediating carcinogenic activities like metastasis activity, tumor cell proliferation, and estrogen-related responses [89]. Melatonin may affect DNA methylation by modulating the activity of DNA methyltransferases involved in the regulation of gene expression by changing DNA methylation patterns. The well-established fact that different tissues present specific patterns of epigenetic modifications [118] may account for the observed tissue-specific effects of ALAN and melatonin on DNA methyl-transferase activity and GDM levels. Tissue differential effects on the activity of DNA methyl-transferases and GDM levels in response to ALAN exposure may present tissue-specific responses to genes that are involved in circadian regulation of several transduction pathways including cancer cell proliferation and metastatic activity. Since humans are diurnal species and most studies have been conducted on nocturnal animals, a diurnal experimental model should be of a great clinical interest. P. obesus may be very useful as a diurnal animal model for understanding the physiological and molecular effects of light pollution on public health. Melatonin suppression, GDM, and even thyroxin levels may present a significant clinical importance as a biomarker for early detection of cancer, particularly in individuals who are at increased risk of developing cancer by circadian disruption induced by excessive ALAN exposures. As epigenetic modifications are revisable, these biomarkers retain therapeutic value.
for ALAN-induced cancer by gene demethylation. Finally, the accumulating data regarding the adverse effects of light pollution on ecology and health compel us to take drastic and rapid measures to reduce light pollution by extreme regulation or at least reducing SWL emission by developing safe lightning technology.
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Dedication

This chapter is dedicated to the memory of Professor Abraham Haim, who passed away before publication of this work. His contribution was foremost among the authors of this chapter.

Abbreviations

ALAN  artificial light at night  
AA-NAT  aryl-alkyl-amine-N-acetyltransferase  
Ws  body mass  
GDM  global DNA methylation  
HIOMT  hydroxyindole-O-methyltransferase  
SCN  hypothalamic suprachiasmatic nucleus  
HPA  hypothalamic-pituitary-adrenal  
HPG  hypothalamic-pituitary-gonadal  
HPT  hypothalamic-pituitary-thyroid axis  
ipRGCs  intrinsically photosensitive retinal ganglion cells  
LED  light-emitting diodes  
RHT  retinohypothalamic tract  
SWLs  short wavelengths  
SCG  superior cervical ganglion  
TSH  thyroid-stimulating hormone

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