INTRODUCTION

Cycles of sleep and wake occur within a circadian chronobiological context. Exposure to sunlight during the day and darkness at night optimally entrains sleep-wake and other biological rhythms to promote homeostasis and human health. Environmental light synchronizes the master circadian pacemaker in the suprachiasmatic nuclei (SCN) of the hypothalamus, as well as many peripheral clocks in tissues and cells, to the solar 24-hour day. Circadian sleep disruption is common in industrialized societies as a result of night shift work, repeated transmeridian flights and mistimed sleep and has been linked to a number of pathological conditions, including hormone-dependent cancers.

On the basis of limited human evidence and sufficient evidence in experimental animals, in 2007 a Working Group from the International Agency for Research on Cancer (IARC) based in Lyon, France classified ‘shift work that involves circadian disruption’ as ‘probably carcinogenic to humans,’ Group 2A [1]. In other words, the IARC which is part of the World Health Organization (WHO), placed shift work in the same category as occupational exposures in petroleum refining and chronic infection with hepatitis C virus in terms of cancer risk (Table 1) [2]. Thus, in 2009, Denmark has become the first country to pay government compensation to women who developed breast cancer after long spells of working at night [3].

Recently, in June, 2019, using the new and updated data, the IARC finalized its evaluation of the carcinogenicity of shift work. Again, the IARC Working Group classified shift work in Group 2A, ‘probably carcinogenic to humans,’ based on limited evidence of cancer in humans and sufficient evidence of cancer as well as strong mechanistic evidence in experimental animals [4].

This review aims to enable a better understanding of the complex relationship between circadian sleep distribution and cancer risk. We first briefly present possible cellular and neuro-endocrine mechanisms involved in this relationship. We then highlight epidemiological data supporting the potential for circadian sleep distribution to act as a carcinogenic agent in the development of hormone-dependent breast and prostate cancers. Finally we address some precautionary measures to limit the risk of carcinogenesis in populations exposed to light at night (LAN).
Circadian Sleep Disruption and Cancer Risk

Regulate cell proliferation and apoptosis at multiple sites. Dysregulation in clock gene expression is associated with increased risk of breast cancer, especially among younger women [7,8]. In addition, variations in core circadian gene cryptochrome 2 (CRY2) is known to increase both risk of developing non-Hodgkin’s lymphoma and prostate cancer risk [9]. In contrast, some polymorphisms in Npas2 appeared to be protective against these malignancies [10].

Nocturnal melatonin suppression by light as a cancer risk factor

Melatonin produced by pineal gland at night is the biochemical correlate of darkness and may impact initiation, promotion and progression of tumor growth. Chronic exposure to artificial LAN results in a significant decrease in melatonin release, and appears to cause tumor development in endocrine target tissues at least by two mechanisms including dysregulation in sexual hormones synthesis and increase in oxidative stress at the cellular level [11].

In experimental models, suppression of melatonin via constant light exposure or pinealectomy augments levels of estrogens and androgens and contributes to increase the incidence of breast and prostate malignancies in a dose dependent manner [11]. In contrast, increased melatonin levels have been shown to inhibit or to slow down tumor growth both in vitro and in vivo, including breast [12,13] and prostate cancers [14].

Melatonin, an effective antioxidant may limit the initiation of cancer by protecting cells from severe DNA damage that is consequence of unstable oxygen-derived and nitrogen-derived toxic reactants (free radicals) [15]. Melatonin and its metabolites act both directly by scavenging free-radicals [15] and indirectly by stimulating antioxidative enzymes [16], which metabolize the toxic compounds to innocuous metabolites. In addition to protecting DNA from destructive oxidant carcinogens, melatonin may also enhance DNA repair capacity by affecting several key genes involved in DNA damage responsive pathways [17].

Dysregulation of circadian cortisol rhythm as a cancer risk factor

Cortisol level is regulated via the hypothalamic-pituitary-adrenal (HPA) axis and shows a strong circadian rhythm with a peak in the morning, 30–45 min after first awakening and a nadir during sleep [18]. Insults to the circadian system (i.e., chronic sleep deprivation and shift work) may alter the response of the HPA axis to the circadian cues responsible for the physiologic pattern of the diurnal cortisol curve [19]. Poorly coordinated cortisol levels with permanent flattened diurnal profile and circadian rest-activity rhythms have been linked with advancing cancer and accelerated tumor growth rates [20,21] as well as increased risk of mortality in metastatic breast cancers [22].

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**Table 1. IARC–WHO identification of carcinogenic hazards to humans**

| Group | Grade | Agents, examples |
|-------|-------|------------------|
| 1     | Carcinogenic to humans | 120 agents: arsenic and inorganic arsenic compounds, painter (occupational exposure as a), estrogen-progestogen oral contraceptives (combined) |
| 2A    | Probably carcinogenic to humans | 82 agents: night shift work, petroleum refining (occupational exposures in), non-arsenical insecticides (occupational exposures in spraying and application of), hepatitis C virus (chronic infection with) |
| 2B    | Possibly carcinogenic to humans | 311 agents: printing processes (occupational exposures in), textile manufacturing industry (working in), pickled vegetables (traditional Asian) |
| 3     | Not classifiable as to its carcinogenicity to humans | 500 agents: silicone breast implants, bisphenol A diglycidyl ether (araldite), triethylene glycol diglycidyl ether |

Based on the evidence available, the IARC–WHO grades potential cancer-causing hazards, and then groups them accordingly. National health agencies can use this information as scientific support for their actions to prevent exposure to potential carcinogens.

IARC: International Agency for Research on Cancer, WHO: World Health Organization.

Adapted from https://monographs.iarc.fr/agents-classified-by-the-iarc/ [2].

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**CELLULAR AND NEURO-ENDOCRINE MECHANISMS OF CIRCADIAN DISRUPTION AND CARCINOGENESIS**

The dominant environmental factor that can reset and thereby disrupt the circadian rhythm is LAN [5]. Chronic exposure to LAN and behavioral disruptions to circadian rhythms (i.e., repeated transmeridian flights, night or rotating shift work, or mistimed sleep) result in a desynchronisation of the master circadian pacemaker with many self-sustained and autonomous peripheral oscillators in tissues and cells [6].

This internal desynchrony between the master circadian pacemaker in the SCN and peripheral oscillators is hypothesized to increase cancer risk through at least three related pathways: 1) direct disruption of the circadian clock genes function that control cell proliferation, 2) nocturnal suppression of the pineal hormone melatonin, and 3) dysregulation of circadian cortisol rhythm.

**Circadian clock genes and carcinogenesis**

Around a dozen of circadian clock genes (peripheral oscillators) regulate cell proliferation and apoptosis at multiple sites. Dysregulation in clock gene expression is associated with the risk of developing cancers by impacting on the biological pathways that regulate DNA damage and repair, carcinogen metabolism, cell-cycle and apoptosis [6]. Structural variations in the circadian gene PER3 and non-synonymous polymorphisms in the circadian gene NPAS2 were detected to be significantly associated with increased risk of breast cancer, especially among younger women [7,8]. In addition, variations in core circadian gene cryptochrome 2 (CRY2) is known to increase both risk of developing non-Hodgkin’s lymphoma and prostate cancer risk [9]. In contrast, some polymorphisms in Npas2 appeared to be protective against these malignancies [10].

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Cortisol would act by stimulating the P450 aromatase activity, which is involved in the conversion of adrenal androgens to estrogens in adipose tissue and muscle [23,24]. It has been suggested that increased aromatase activity leads to increased breast and prostate cancer risks [23,25]. Due to the natural involvement of cortisol in the functioning of the prostate and mammary glands, its circadian dysregulation and prolonged diurnal presence in blood and tissues is likely to expose prostate and breast cells to the activation of downstream biological pathways outside their normal context [26].

EPIDEMIOLOGICAL EVIDENCES

Epidemiological data on hormone-dependent breast and prostate cancer risks linked to disturbances of the circadian sleep rhythm can be analyzed from both night shift work and habitual sleep duration perspectives.

Breast cancer

Breast cancer is the most commonly occurring cancer in women worldwide, with the highest rates found in the Western Europe, Australia and North America [27]. Breast cancer risk is affected by a combination of several factors including genetic predisposition, advanced age, lifetime exposure of breast tissue to hormones (early menarche, late menopause, and hormone use) and lifestyle choices [28]. Modern lifestyle and consequent exposure to artificial LAN also have been reported as a possible risk factor for breast cancer [29,30]. Furthermore, migrant studies have shown that, over successive generations, incidence rates increase in women who migrate from low-incidence countries in Asia and Latin America to high-incidence countries such as the United States and Australia, approaching the incidence rates observed in the host country [31-33].

Shift work and breast cancer

The association between an increased risk of breast cancer and night shift work has been reported by numerous epidemiologic studies, including cohort studies, case-control studies, and meta-analysis [34-40]. The risk of breast cancer seems to be linked to the nature and exposure metrics of night shifts. For some authors, regardless of years of exposition, there is some evidence that high number of consecutive night shifts has impact on the extent of circadian disruption, and thereby increased breast cancer risk. For others, there is a dose-response relationship between years of night shift work exposition and breast cancer risk. In a cohort study of 102,869 women aged 16 or older from the UK during a 9.5-year median follow-up period, there was a significant link between raised breast cancer risk and average hours of night work per week (p=0.035) [40]. A Norwegian cohort of 43,316 nurses has also reported a significant increase in the risk among nurses who worked at least 5 years performing at least 6 consecutive night shifts per month [odds ratio (OR): 1.8, 95% confidence interval (CI): 1.1–2.8] [38]. These results suggest that the risk of breast cancer may be related to number of consecutive night shifts. A dose-response relationship between rotating night shift work and breast cancer has emerged in the US based Nurse Health Study [34]. This prospective cohort involved 78,862 nurses who had worked at least three nights per month and followed over a 10-year period. Authors have reported a significant increase in breast cancer risk among the women who worked 30 or more years on rotating night shifts indicating a multivariate adjusted relative risk (RR) of 1.36 (95% CI: 1.04–1.78) [34]. In an additional study, same authors followed 115,022 nurses over a period of 12 years. Breast cancer was diagnosed in 1,352 of them. The RR for breast cancer was 1.79 (95% CI: 1.06–3.01) for women who worked at least 3 night shifts per month over a period of 20 years [35]. In contrast, a meta-analysis including 10 prospective studies concluded that night shift work for 20 or more years has little or no effect on breast cancer incidence [41]. However, several shortcomings with this meta-analysis have been underlined: 1) exclusion of all retrospective studies as uninformative, 2) inclusion of studies where the assessment of shift work fell below the IARC working group standards, 3) assessment of shift work often based on self-report (and so was not collected prospectively), and 4) the heterogeneity in definitions of shift work that contraindicate the data being combined in a meta-analysis [42]. Based on a comparative case-control study of 1,679 Israeli women, exposure to LAN in the “sleeping habitat” has been found to be associated with breast cancer risk (OR: 1.22, 95% CI: 1.11–1.31; p<0.001) [43]. Authors concluded that, not only should artificial light exposure in the working environment be considered as a potential risk factor for breast cancer, but also LAN in the “sleeping habitat.”

Sleep duration and breast cancer

Several observational studies have investigated the association between sleep duration and breast cancer risk with conflicting results: 1) short sleep duration might increase the risk and longer sleep duration might decrease this risk, 2) longer sleep duration might increase the risk, 3) sleep duration has no impact on the risk, 4) sleep duration might decrease this risk, 2) longer sleep duration

First, a significant number of population-based observational studies reported that short sleep duration is associated with an increased risk [44,45], or aggressiveness [46], and longer sleep duration with a decreased risk of breast cancer [47,48]. In the Ohsaki Cohort Study of 23,995 Japanese women, participants who slept 6 hours or less compared to a reference group of women who slept 7 hours were more likely to have a breast cancer with a multivariate hazard ratio (HR) of 1.62 (95% CI: 1.05–2.50; p for trend= 0.03) [44]. Later, the Southern Community Cohort Study carried out to investigate the role of sleep duration in breast cancer among African American women found results similar to those observed in Japanese females. Compared to the reference group with an average sleep of 8 hours per day, black women with shorter sleep had an increased risk of hormone receptor negative breast cancer [ORs (95% CIs): 2.13 (1.15–3.93), 1.66 (0.92–3.02) and 2.22 (1.19–4.12) for <6, 6 and 7 hours, respectively, p for trend, 0.04]
were 1.10 (95% CI, 0.59–2.05), 1.0, and 0.28 (95% CI, 0.09–0.88), respectively. A significantly lower risk (p=0.03) in women who reported a long duration of sleep (≥9 hours per night) compared with the average duration of 7 or 8 hours was observed. No statistically significant effects were observed for sleep insufficiency and sleep quality. Wu et al. [48] investigated the relationship between self-reported usual sleep duration determined at baseline and subsequent risk of breast cancer in the prospective, population-based cohort of the Singapore Chinese Health Study. During the follow-up period (up to 11 years), 525 incident breast cancer were identified among 33,528 women participating in the study. Among women postmenopausal at baseline, breast cancer risk decreased with increasing sleep duration (p=0.047). Women who reported ≥9 hours of sleep per night showed an RR of 0.67 (95% CI: 0.4–1.1) compared with those who reported ≤6 hours of sleep per night.

Second, only a few studies have stated that longer sleep duration may increase the risk of breast cancer. One case-control study from the US [49], including premenopausal and postmenopausal breast cancers, found that self-reported longer sleep duration in the past 2 years slightly increased the risk of breast cancer (9 or more versus 7–8 hours of sleep per night). The multivariate-adjusted OR for developing breast cancer was 1.06 (95% CI: 1.01–1.11), suggesting a 6% increase in risk for every additional hour of sleep. According to a more recent meta-analysis based on 10 studies involving 415,865 participants; compared to women with the reference number of sleep hours (6 to 7 hours), women with a longer sleep duration might have a significantly increased risk of breast cancer, especially estrogen receptor-positive breast cancer (p for trend=0.024) [50].

Third, in some other studies, no association was observed for shorter or longer sleep duration. Vogtmann et al. [51] investigated the Women's Health Initiative (WHI) consisted of four clinical trials and one observational study to determine whether the duration of sleep, sleep quality, insomnia, or sleep disturbance was associated with incident breast cancer. A total of 5,149 incident cases of breast cancer were identified in this study. No statistically significant associations were found between self-reported sleep duration, sleep quality, insomnia, or level of sleep disturbance with the risk of breast cancer after multivariable adjustment. Pinheiro et al. [52], explored the association between habitual sleep duration and risk of breast cancer in the Nurses’ Health Study including 77,418 women. During 16 years of follow-up, 4,223 incident cases of breast cancer occurred. Compared with women sleeping 7 hours, covariate-adjusted HRs and 95% CIs for those sleeping 5 or less, 6, 8, and 9 or more hours were 0.93 (0.79–1.09), 0.98 (0.91–1.06), 1.05 (0.97–1.13), and 0.95 (0.82–1.11), respectively. In conclusion, authors found no convincing evidence for an association between self-reported sleep duration and the incidence of breast cancer.

Fourth, a population-based case-control study from Guangzhou, China indicates a U shaped relationship between sleep duration and breast cancer [53]. Wang et al. [53] conducted face-to-face interviews with 712 women diagnosed with incident invasive breast cancer before treatment and 742 age-matched controls. Among all subjects, 33.0% of cases and 26.2% of controls reported ever having night-shift work [OR (95% CI): 1.34 (1.05–1.72)]. Compared to women with an average sleep duration (6.1–8.9 hours per day), women who had shorter (≤6.0 hours per day) [OR (95% CI): 1.53 (1.10–2.12)] and longer sleep duration (≥9.0 hours per day) [OR (95% CI): 1.59 (1.17–2.17)] had an increased risk of breast cancer. Night-shift work was associated with an increased risk of breast cancer [OR (95% CI): 1.34 (1.05–1.72)]. In addition, daytime napping was associated with a reduced risk of breast cancer among night-shift workers [OR (95% CI): 0.57 (0.36–0.90), but no association was found among women who never had night-shift work. Night-shift work and longer sleep duration also synergistically increased breast cancer risk [OR (95% CI): 3.69 (1.94–7.02)] (p for interaction=0.009). Sleep problems, including night-shift work, and shorter and longer sleep duration, are associated with an increased breast cancer risk. In particular, the combined effects of night-shift work with no daytime napping or longer sleep duration are greater than the independent effects.

Prostate cancer
Prostate cancer is one of the most common cancers in males with an increasing incidence worldwide [54]. The etiology of this hormone-dependent tumor seems multifactorial and remains largely unknown compared to other common cancers. Chronic sleep deprivation and night shift work have been suggested as possible risk factors for prostate cancer [55].

Shift work and prostate cancer
Epidemiological studies on shift work and prostate cancer risk
are scarce and results are mixed. A meta-analysis including 2,459,845 individuals from eight studies published between 2006 and 2014 (three case-control studies and five cohorts) found a 24% increase in the risk of prostate cancer for night-shift workers (RR: 1.24, 95% CI: 1.05–1.46; p=0.011) [56]. Dose-response subgroup analysis suggested that an increase in night-shift work of 5 years duration was significantly associated with a 2.8% (95% CI: 0.3–5.4%, p=0.030) increase in the risk of prostate cancer.

A more recent French population-based case-control study [57], including 818 incident prostate cancer cases and 875 controls reported that overall, ever night work, either permanent or rotating, was not associated to prostate cancer. Similarly, a prospective cohort study of Finnish twins [58] including 11,370 men, over a median of 30 years of follow-up did not find any association neither with rotating shift work (HR: 1; 95% CI: 0.70–1.20), nor with fixed night work (HR: 0.5; 95% CI: 0.10–1.90). However, according to the dose-response analysis in the French study [57], a long duration of at least 20 years of permanent night work was associated with aggressive prostate cancer (OR: 1.76, 95% CI: 1.13–2.75), even more pronounced in combination with a long shift length (>10 hours) or at least 6 consecutive nights (OR: 4.64, 95% CI: 1.78–12.13; OR: 2.43, 95% CI: 1.32–4.47, respectively).

**Sleep duration and prostate cancer**

In a population-based Swedish cohort study (n=14,041 men), during 13 years of follow-up, Markt et al. [59] identified 785 cases of incident prostate cancer. Overall, 20% of men reported sleeping 8 hours per night, 4% reported sleeping 5 hours or fewer, and 2% reported 9 hours or more of sleep per night. Men at the extremes of sleep duration (≤5 h and ≥9 h) were more likely to be retired and therefore have no working hours. In multivariable-adjusted analyses, sleep duration was not associated with risk of prostate cancer. In addition, there were no association between prostate cancer and sleep disruption, as defined by difficulty falling asleep, difficulty maintaining sleep, sleep quality, and restorative power of sleep.

In contrast, two other studies indicate that, short night-time sleep may be associated with an increased risk of prostate cancer in non-shift working men. Within the prospective AGES-Reykjavik population-based cohort (n=2,102 men), the association between sleep disruption and prostate cancer risk has been studied over a five-year time frame [60]. During follow-up, 135 men (6.4%) were diagnosed with prostate cancer. Compared to men without sleep disruption, those with problems falling and staying asleep were at significantly increased risk of prostate cancer [HR, 1.7 (95% CI: 1.0–2.9) and 2.1 (95% CI: 1.2–3.7)], respectively, with increasing sleep disruption severity. In other words, men with insomnia were twice as likely to develop prostate cancer. In the Ohsaki Cohort Study [61], an inverse association between sleep duration and the risk of prostate cancer has been found among non-shift working Japanese men. Authors reported a decreased risk for those sleeping at least 9 hours per night. The HR of prostate cancer in short sleepers (those sleeping 6 hours or less) was 1.34 (95% CI: 0.83–2.17) and the HR for long sleepers (those sleeping at least 9 hours) was 0.48 (95% CI: 0.29–0.79) (p for trend=0.02).

**CANCER RISK MANAGEMENT IN THE CONTEXT OF CIRCADIAN SLEEP DISRUPTION**

In the globalizing world, many countries face to the growing demands in productivity in different industries such as energy production, transportation, health care, telecommunication and entertainment. Consequently, circadian sleep disruption related to modern life conditions including the night-time use of portable light-emitting devices (i.e., computer screens, tablets and cell phones) [62] and extended nocturnal business hours and night shift works are increasingly prevalent in the general population and even a modestly increased risk could lead to considerable numbers of breast and prostate cancer cases if there were a causal association.

While many factors and substances in living and working environments are known to be carcinogenic, others are suspected to be carcinogenic such as circadian sleep disruption [1]. In the area of risk management, it is important to distinguish prevention, which is the management of known and proven risks, from precaution, which corresponds to the management of uncertain risks. The precautionary principle is based on the use of public management of potentially serious and/or irreversible health risks, especially when there are significant uncertainties regarding the impact on the population. The precautionary principle is usually associated with the principle of proportionality, according to which the cost of the measures necessary to reduce the risk must not be disproportionate to the expected benefits. Thus, a number of precautionary measures may be proposed in order to reduce the possible impact of chronic circadian sleep disruption on the genesis of breast and prostate cancers. The years of night shifts can be limited for employees (i.e., less than 20 years), the duration of night shift working hours may be shortened. High number of consecutive night shifts may be avoided (i.e., fewer than 6 consecutive nights per month). Instead of permanent night work, rotating shifts may be proposed. Rest periods after night shifts may be lengthening. Finally, co-exposure to other occupational carcinogens additional to years of employment in non-day shift work may be controlled. In addition, melatonin with its oncostatic properties may be of use in the precautionary measures and treatment of breast and prostate cancers [11].

**CONCLUSION**

This review illustrates the emerging evidence from human and animal studies that chronic circadian sleep disruption related to sleep deprivation and/or night shift work may be a significant risk factor for both breast and prostate cancers. The mechanistic hypotheses that the multilevel endocrine changes caused by circadian disruption with sleep deprivation and melatonin suppression...
through LAN may favor the induction and/or promotion of malignant tumors are likely [63]. Despite a large number of epidemiological studies suggesting that circadian sleep disruption due to night shift work may be associated with increased risk of both breast and prostate cancers, some other cohorts and meta-analyses did not find any associations. These conflicting results might be due to several methodological limitations. First, in available epidemiological studies there have not been clear and uniform definitions of ‘shift work’ and ‘shift work exposure times’ used [1]. Second, the majority of these studies did not take daytime napping into account [53] as well as the possible environmental carcinogenic factors associated to sleep deprivation and night shift work. Third, short-term follow-up of aging cohorts may explain why some cohort studies may have null findings [64]. Consequently, the need for high quality epidemiological cancer research still exists. In the future, epidemiological studies might be designed with more precise definitions of shift work, sleep deprivation and nocturnal light exposure. Genetic profiles (polymorphisms in circadian genes, clinical chronotypes), objective sleep patterns (actimetric and polysomnographic findings), other environmental and occupational carcinogenic factors and comorbidities can be integrated into analyses. Since the classification of shiftwork that involves circadian disruption as a ‘probable human carcinogen’ by the IARC, subjects who have worked 20 years or more in night shift work may need special attention for the screening and monitoring of breast and prostate cancers. Finally, based on the precautionary principle, at-risk subjects can be identified and moved away from shift work stations as early as possible before reaching several years of exposure. In order to limit the risk of cancer, this precautionary principle needs to be applied to public health actions as well as to actions pursued by policy makers, and industry.

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Conflicts of Interest
The authors have no potential conflicts of interest to disclose.

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