The inner clock—Blue light sets the human rhythm

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Abstract
Visible light synchronizes the human biological clock in the suprachiasmatic nuclei of the hypothalamus to the solar 24-hour cycle. Short wavelengths, perceived as blue color, are the strongest synchronizing agent for the circadian system that keeps most biological and psychological rhythms internally synchronized. Circadian rhythm is important for optimum function of organisms and circadian sleep–wake disruptions or chronic misalignment often may lead to psychiatric and neurodegenerative illness. The beneficial effect on circadian synchronization, sleep quality, mood, and cognitive performance depends not only on the light spectral composition but also on the timing of exposure and its intensity. Exposure to blue light during the day is important to suppress melatonin secretion, the hormone that is produced by the pineal gland and plays crucial role in circadian rhythm entrainment. While the exposure to blue is important for keeping organism’s wellbeing, alertness, and cognitive performance during the day, chronic exposure to low-intensity blue light directly before bedtime, may have serious implications on sleep quality, circadian phase and cycle durations. This rises inevitably the need for solutions to improve wellbeing, alertness, and cognitive performance in today’s modern society where exposure to blue light emitting devices is ever increasing.

KEYWORDS
visible light, blue light, circadian rhythm, melanopsin, melatonin

1 | THE ADVENT OF THE CIRCADIAN PHOTORECEPTOR

Circadian rhythms are biological cycles that have about 24-hour periods. Body temperature, hormonal levels, sleep duration and quality, cognitive performance and countless other physiological variables exhibit such daily oscillations. The ability of the visible light to synchronize the biological clock of the photosynthetic marine dinoflagellate Gonyaulax polyedra to the solar 24-hour cycle is known since 1958 [1]. It was not until 1998, when a new type of photoreceptor in the human eye was discovered [2] and later identified to be especially sensitive to blue light. Importantly, these new photoreceptors are retinal ganglion cells (RGCs) and communicate directly with the brain. The melanopsin containing retinal ganglion cells, called intrinsic photosensitive retinal
ganglion cells (ipRGCs) count for around 1% of the total RGCs and are found in front of the retina (see Figure 1) to process and send signals [3–5] among others to the suprachiasmatic nucleus (SCN), the so called master clock of the brain [6]. In depth morphological analyses allowed for an ipRGC population clustering into five distinct subtypes, which moreover express a certain diversity in their dendritic stratification, morphology, membrane properties, and melanopsin expression levels [7]. Worth mentioning are hereby the most light sensitive M1 cells, which in turn can be separated according to their Brn3b expression, with Brn3b-negative cells operating as the herein described reporters to the SCN, but likewise signal to the olivary pretectal nucleus to influence the pupillary light reflex [7]. Other, less light sensitive, subtypes project to further, even image forming, brain areas, indicating potentially different functional responsibilities, thereby influencing many aspects of human physiology [7]. These circadian photoreception cells differ from the classical rod and cone photoreceptors in many ways. They use a unique photopigment, melanopsin and have lower sensitivity and spatiotemporal resolution than rods or cones. Light depolarizes ipRGCs but hyperpolarizes rods and cones. Further, ipRGCs are less sensitive than the classical photoreceptors and are far more sluggish, with response latencies as long as 1 minute. Their strongly overlapping dendrites, which are themselves filled with melanopsin, are able to respond directly to light [4], which is why the entire ipRGC population has been referred to as” photoreceptive net”[8]. Bright continuous illumination evokes a sustained depolarization in ipRGCs that encodes stimulus energy. This sets these cells apart from all other photoreceptors, which are less sensitive to encode irradiance and thus to represent ambient light intensity [5, 9, 10] over days [11] and months [12]. These properties are well suited to nondirectional detection of gross environmental illumination essential for integrated circadian, neuroendocrine and neurobehavioral effects [5]. They also contribute to the pupillary light reflex and other behavioral and physiological responses to environmental illumination. The M1 Brn3b-negative ipRGC photoreceptors send unconscious, nonvisual photic information through the retinohypothalamic tract to the SCN (see Figure 2) permitting alignment of internal biological with external environmental time. Cells in the SCN thereby create a circadian rhythm via interconnected gene-protein feedback loops, wherein transcription and activity of clock genes and transcribed proteins function as their own regulators [13]. Worth mentioning is the control loop including Period (Per1, Per2, and Per3) and Cryptochrome (Cry1 and Cry2) genes, whose transcription is promoted by transcription factors Circadian Locomotor Output Cycles Kaput (CLOCK) and Brain and Muscle ARNT-like protein 1 (Bmal1), whose transcribed proteins Per & Cry in turn inhibit CLOCK/Bmal1-mediated transcription [13]. These
feedback-loops in turn require external cues, among others light [4], social interactions [14] and nutrition [15], to maintain a 24-hour schedule [13]. The resulting SCN pacemaker neuron activity subsequently synchronizes the rhythms in subsidiary clocks of peripheral tissue, which would otherwise run free and desynchronize [16]. Therefore, although absent or deficient ipRGC photoreception cannot be perceived subjectively [17], the missing entrainment yields in circadian disturbances, which can have significant physiological and psychological consequences [18].

2 | IMPORTANCE OF BLUE LIGHT IN REGULATING HUMAN BEHAVIOR AND CIRCADIAN RHYTHM

Cones, the color receptors of the human visual system, are classified according to the wavelength sensitivity peak of their respective photopigments, wherefore they are commonly referred to as short (S, 420 nm), medium (M, 533 nm), and long (L, 562 nm) wavelength cones [19]. Under photopic conditions, these spectral opsin sensitivities give rise to the photopic luminosity function, describing the relative sensitivity of the human eye to white light intensities, with a peak around 555 nm [20]. Rods on the other hand only rely on one photo pigment, rhodopsin, which is most sensitive at 498 nm [19]. Due to their retinal ganglion cell nature, ipRGCs receive, besides their own melanopsin based photosensitivity, intraretinal synaptic input from classical image-forming photoreceptor driven circuits [21]. This input potentially modulates the nonimage-forming ipRGC light response transmitted to higher brain regions [21]. The extrinsic photoreceptor influence is most notably light intensity dependent, whereby rods impact the intrinsic ipRGC response primarily in scotopic conditions (~7 log photons cm$^{-2}$ s$^{-1}$), while cones contribute in photopic conditions (~11 log photons cm$^{-2}$ s$^{-1}$), although less pronounced [22, 23].

Taking this photoreceptor intensity sensitivity, the melanopsin phototransduction threshold (~9 log photons cm$^{-2}$ s$^{-1}$) and the respective dynamic ranges into account, an ipRGC is able to encode light intensities in a range of 9 orders of magnitude, with modestly varying spectral sensitivities as a results of extrinsic photoreceptor input [24]. Sensitivity of human nocturnal melatonin suppression [25, 26] peaks in the blue portion of the light spectrum around 460 nm, most effective in the range between 446 and 477 nm, manifesting a substantial contribution of short wavelength visible light not directly corresponding with an established single opsin maximum sensitivity [27]. Spectral absorption by melanopsin [28], which is the photopigment expressed in cell bodies and elongated dendrites [5] of ipRGCs, peaks at 479 nm [29]. The difference of ~20 nm might reflect an influence of short wavelength cones on the melatonin response. Thus, blue wavelengths can potentially exert more powerful effects on human circadian rhythm than green and yellow wavelengths. This possibility has been confirmed experimentally [30] when suppression of melatonin in humans was compared during nighttime exposure for monochromatic light at 460 nm with light at 555 nm, the peak sensitivity of the visual cones. The blue wavelength suppressed melatonin for about twice as long as green, despite the equal photon density. Blue monochromatic light exposure was more effective to cause a phase delay of the circadian rhythm of all subjects, which were studied in an environment free of time cues. Circadian phase assessments were done by monitoring the melatonin secretion profiles (see Figure 3 for a physiological profile), before and after monochromatic light exposure (6.5 hours), while subjects performed constant routines. While the photon flux was matched for different wavelengths, the observed effect that blue light was more effective in circadian shifts suggests that ipRGCs do not simply count or average photons but are rather sensitive to the particular wavelength of exposure [30]. Similar wavelengths also proved more powerful in elevating body temperature and heart rate and in reducing sleepiness, while provoking responses in alertness-related subcortical structures and limbic areas, which established light, most notably in the blue range, as a potent modulator of brain function [31, 32].

Evidence for the importance of light in regulating the circadian rhythm also come from studies that used bright light to treat mood disorders [34]. Melatonin suppression in humans was observed by applying bright light of 2500 lx, much brighter than indoor lighting, but much less so than a cloudy day. Therefore, it was concluded that the circadian rhythm is best cued by natural bright light and would be insensitive to dim indoor lighting. The finding that bright light can suppress melatonin [34] led to treating seasonal affective disorder with bright light. The literature on the efficacy of blue light for treating this disorder is just now beginning to develop, but research on the efficacy of blue light to suppress melatonin suggests that exposure to blue light might be more effective treatment. Blue light may further prove useful for treating premenstrual depression, bulimia, and anxiety [32]. Personalized blue light exposure raised activity levels during daytime hours and increasing sleep at nighttime in a long-term care facility for patients with Alzheimer’s disease and related dementia [35]. An investigation of dysfunction-pathologic structure relationships of the hypothalamus in Alzheimer’s disease patients could likewise prove a direct effect of light therapy on circadian rhythm related disturbances [36].

Further, a correlation of polychromatic light intensity and nocturnal melatonin suppression could be demonstrated,
with 1 hour of 1000 lx exposure by night being sufficient to reduce melatonin levels to a daytime average [37]. Studies indicate that low levels of illumination are sufficient to trigger awakening and there are hints that the effect on circadian rhythm gets saturated at high illumination levels [38, 39]. Electroencephalography has shown that light exposure influences alpha, theta, and low-frequency activity, which are correlates of sleepiness [31]. Blue light proved superior to other wavelengths in enhancing responses in the left frontal and parietal cortices during a working memory task. Experimental subjects had quicker auditory reaction times and fewer lapses of attention under blue light than green light [31]. Blue wavelengths suppressed sleep-associated delta brainwaves better than green wavelengths and boosted the alpha wavelengths, which are related to alertness [31]. This suggests that short wavelengths, perceived as blue color, might be used to control sleepiness. Therefore, blue light seems to be the strongest synchronizing agent for the circadian system that keeps most biological and psychological rhythms internally synchronized. The sleep/wake cycle is however not to be equated with circadian rhythms, since additional factors, most notably the level of homeostatic sleep drive, which only diminishes by sleep itself, affect sleep propensity. The expression of circadian rhythms nevertheless seems to be most obvious in the rhythmicity of sleep and neurobehavioral performance in humans [40]. It has further been hypothesized that light has additional, non-circadian effects on alertness and the sleep/wake cycle, most notably by acting on sleep promoting, as the ventrolateral preoptic nucleus, and wake promoting brain regions either directly or eventually via the SCN [41]. These conclusions reached further, as to suggest, that rod and cone encoded light stimuli were primarily involved in the promotion of the wake state, whereas melanopsin primarily evokes a higher level of alertness by mediating EEG correlates of waking associated cognitive processes [41]. It is recognized, that subjective alertness is moreover influenced by subjective aspects, such as well-being and visual comfort [42]. The power to reset the organism's daily meter declines into green (505 nm) [43] and green-yellow (555 nm) colors [30]. Thus, today it is understood that blue light has many physiologic effects but perhaps the most important among them is to entrain the circadian rhythm [5]. A selection of studies, investigating the effects and details of spectral composition, intensity, duration, timing and temporal distribution of light stimuli [44] on human behavior and circadian rhythm are outlined in Table 1 of the appendix.

3 | BLUE LIGHT IMPACT ON AGING AND RETINAL DAMAGE

Age-related changes in lens density [45, 46] are known to reduce the transmission of blue light, which has been shown to be most effective in suppressing melatonin secretion during the day. A further age-related change in the eye that may contribute to reduced levels of light reaching the ipRGCs is the reduction in pupil size [47]. Therefore, this diminished
TABLE 1  Selective overview of published studies and methodological parameters regarding the effect of light, most notably in the blue spectrum, and illumination intensity on circadian rhythms, alertness, and sleep. Most studies in humans included an initial ophthalmological examination and evaluation of chronotype, constant posture protocols in the laboratory and dark adaptation episodes under polychromatic dim light for 0.5 to 2 hours before treatment exposure. Irradiance unit summarizes information about photon density, irradiance, illuminance, and luminance. The properties timing and duration refer to start and length of the respective treatment exposure, not the whole experimental protocol.

| Study                  | Methodology                             | Spectral parameter | Unit                             | Duration | Timing            | Subjects (mean ± SD) | Results                                                                 |
|------------------------|-----------------------------------------|--------------------|----------------------------------|----------|-------------------|----------------------|-------------------------------------------------------------------------|
| Lockley et al [30]     | BMA                                     | 555 and 460 nm     | $6 \times 10^{13}$ photons cm$^{-2}$ s$^{-1}$ | 6.5 h    | 9.25 h before wake time | 16 healthy, (23.3 ± 2.4) | 460 nm induced a 2-fold greater circadian phase delay                   |
| McIntyre et al [37]    | BMA                                     | polychromatic      | -                                | 1 h      | Midnight          | 13 healthy, (25.1 ± 6.4) | 1000 lx intensity suppress melatonin to near daytime levels; 350 lx significantly suppress nocturnal melatonin levels. |
| Munch et al [43]       | Sleep: PSG, 8 EEG, EOG, EMG, ECG         | 550 nm, 460 nm,    | 460 nm: 12.1 μW cm$^{-2}$, 550 nm: 10.05 μW cm$^{-2}$ | 2 h      | 9:30 PM          | 8 healthy, male (24.6 ± 3)  | Small wavelength-dependent effects of light on sleep architecture and EEG. Most likely an acute alerting effect continuing into sleep and/or an immediate phase delay induced by blue light. |
| Herljevic et al [48]   | BMA                                     | 548 and 456 nm     | 7000-12 000 lx vs. 150 lx         | 0.5 h    | 3.5 h pre melatonin peak | 13 pre- and 21 post-menopausal women | Reduced melatonin suppression in elderly subjects following exposure to blue light, likely due to age-related changes in lens density. |
| Sletten et al [49]     | Questionnaires: Alertness, sleepiness, mood, BMA | 548 and 456 nm     | 200-400 lx                        | 2 h      | 8.5 h after DLMO  | 11 young (23.0 ± 2.9), 15 old (65.8y ± 5.0) men. | Subjective alterness, sleepiness and mood response to blue light diminished in elderly compared to young subjects. No age effect in green light exposure. |
| Cajochen et al [57]    | KSQ, CBT and surface skin temperature, ECG, SMA | 550 nm, 460 nm,    | $2.8 \times 10^{13}$ cm$^{-2}$ s$^{-1}$ | 2 h      | 10 h after wake time | 10 healthy, male (25.9 ± 3.8) | Blue light exposure in the evening suppressed melatonin more than green | (Continues)
| Study                        | Methodology                                                                 | Light parameter                        | Subjects (mean age ± SD) | Results                                                                                                                                                                                                 |
|------------------------------|-----------------------------------------------------------------------------|-----------------------------------------|--------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Czeisler et al [59]          | Night shift work treatment; Sleep-wake logs, ECG, CBT, Cognitive-performance tasks | Polychromatic                           | 8 h Midnight             | 8 healthy, male, 22-29 years Maladaptation of the human circadian system to night work can be treated with scheduled exposure to bright light at night and darkness during the day. |
| Davidson et al [62]          | Molecular rhythms, high-resolution optical microscopy & bioluminescence of SCN | Polychromatic                           | 6 h phase shift          | mPer2LUC knock-in mice Differences in circadian shifting kinetics are apparent among subjects and among organs.                                                                                 |
| Khalsa et al [63]            | BMA                                                                         | Polychromatic                           | 6.7 h, alternating gaze  | 21 healthy, entrained Phase delays occur when light stimulus is centered prior to the core body temperature minimum (CBTM), phase advances occur when light stimulus is centered after the CBTM, no phase shift occurs at CBTM. |
| Cajochen et al [68]          | SMA, KSQ, KDT, GO/NOGO task, cognitive performance                         | LED with twice more 464 nm emission    | 5 h 6 h before bed time  | 13 male (23.8 ± 5.0) Evening exposure to blue enriched LED screen resulted in attenuated salivary melatonin and sleepiness levels, accompanied with increase in cognitive performance. |
| Wahnschaffe et al [71]       | Sleep log and actimetry; SMA, VAS                                          | White, zero blue component, high intensity light | 0.5 h 1 h before bedtime | 9 healthy, (26.3 ± 4.2) Yellow light exposure did not alter the increase of melatonin in saliva. Lighting conditions including blue components                                                                 |
| Study | Methodology | Light parameter | Subjects (mean age ± SD) | Duration | Timing | Results |
|-------|-------------|-----------------|--------------------------|----------|--------|---------|
| Chang et al [73] | Blood plasma melatonin assay, PSG, KSQ, EEG | LE-eBook: blue enriched; Print book: white light | 12 healthy, (24.9 ± 2.9) | 4 h | 6 PM | LE-eBook condition compared to print book: suppressed evening levels of melatonin, delayed melatonin onset, delay sleep onset, less rapid eye movement (REM) sleep. |
| Kozaki et al [74] | SMA at 1 AM and 9 AM | Polychromatic white & blue enriched light | 12 healthy, male (21.9 ± 0.9) | 1.5 h | 1 AM, 9 AM | Findings suggest that daytime blue light exposure has an acute preventive impact on light-induced melatonin suppression. |
| Kervezee et al [98] | Night shift protocol: transcriptome assay | Polychromatic mean 2.6 ± 0.4 lx (SD) | 8 healthy | 8 h | 10 h phase shift | Reduction of rhythmic transcripts in the night shift condition. Mainly due to dampened rhythms & lower amplitudes rather than to a complete loss of rhythmicity. |

Abbreviations: BMA, blood melatonin assay; CBT, continuous body core temperature; DMLO, dim light melatonin onset; ECG, electrocardiogram; EEG, electroencephalography; EMG, electromyogramm; EOG, electrooculography; KDT, Karolinska drowsiness test; KSQ, Karolinska sleep questionnaire; PSG, polysomnography; PVT, psychomotor vigilance task; SMA, saliva melatonin assay; VSA, visual analogue scale.
Blue light input to the circadian clock may result in disturbed circadian rhythm and sleep in the elderly. Age-related changes in melatonin suppression were first studied by Herljevic [48], where suppression was compared across light conditions and between age groups. Two groups of young and elderly subjects were exposed for 30 minutes to blue and medium wavelength light at different irradiances. Light exposure was adjusted so that each subject received light on the rising phase of its endogenous melatonin rhythm. Significantly reduced melatonin suppression was measured in the elderly subjects in response to exposure to blue (456 nm) light as compared to the young group. This finding is suggested to reflect the age-related changes in lens density. In another study [49], it was further demonstrated that similar age-related reduction occurs in subjective alertness, sleepiness, and mood following blue light exposure. The reduction in responsiveness to the effects of blue light in older people was not observed following green light exposure. The magnitude of circadian phase advance was assessed by monitoring the difference in plasma melatonin rhythm before and after light exposure [49]. The phase advances to both blue and green light were larger in the young than older subjects, although differences did not reach statistical significance. To analyze how age-related losses in crystalline lens transmittance and pupillary area affect circadian rhythm phakic and pseudophakic individuals of the same age range were studied [50]. In phakic eyes the natural lens is left untouched after intraocular lens (IOL) implantation to correct myopia, while in pseudophakic eyes the natural lens is explanted. The age-related decline in retinal illumination was estimated taking into account human crystalline lens transmittance at different ages and the pupil area for those ages. Pupil-weighted spectral retinal illumination was multiplied wavelength by wavelength with melatonin suppression sensitivity between 350 and 700 nm to determine how aging affects circadian photoreception for different monochromatic lights. Results showed a 10-fold greater circadian photoreception in young children (10 years old) than adults (95 years old) with phakic eyes. Pseudophakia improves circadian photoreception at all ages, particularly with UV-only blocking IOLs that transmit blue wavelengths. A recent study further suggests that the aging of retinal tissue most severely affects the reduction in amplitude of rhythm of the circadian clock [51].

### 3.1 More facets of blue light

While short wavelengths in the range of 460 nm are very efficient in phase shifting of circadian system, intensive blue light in the range between 400 and 440 nm (see Figure 4) is damaging to the retina via a photochemical reaction called photoreversal of bleaching [52]. It augments the capability of rhodopsin molecules to absorb photons by several orders of magnitude, thus allowing the molecules to reach the critical number of photons required to induce damage in the retinal cells. This process can further increase the potential production of reactive oxygen species (ROS). The oxidative damage can lead to the accumulation and build-up of

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**FIGURE 4** Irradiance of the blue light hazard function. The Blue light hazard function according to ISO 8980 represents the relative spectral sensitivity of the human eye to blue light hazards, based on the effectiveness of radiation to induce photoreversal of bleaching. The emission spectrum of an ordinary smartphone screen on maximum brightness shows a distinct overlap in the potentially harmful blue peak area, and melanopsin sensitivity, but also image-forming related S-cone and rod absorbance spectrum. Normalized absorbance spectra are depicted according to Bowmaker and Dartnall, 1980 [19]
lipofuscin in the retinal pigment epithelium (RPE). The build-up of lipofuscin in the RPE can affect its ability to provide nutrients to the photoreceptors, affecting their viability. The severity of the blue-light induced damage depends on the time of the day and is thus related with the circadian rhythm [52]. Accumulation of ROS and lipofuscin may be due to both acute and cumulative exposure to blue light [52]. While the mechanisms behind the effect of long-term sub-threshold exposure to blue light are not fully understood, it seems to be linked to the lipofuscin level in RPE [52]. Lipofuscin is mainly formed in the photoreceptor outer segments as byproduct of the degradation of the disks. When it absorbs blue light, ROS are produced. The type of ROS created is related to the specific wavelength (\( \lambda \)) of the blue light. The amount of lipofuscin in the retina increases with age and is related to age-related macular degeneration [52]. However, the detrimental effect of blue light on the retina should be taken with caution. Most of the damaging effects through blue light are demonstrated in nocturnal animals only, or in isolated animal cell cultures directly exposed to blue light [52–54]. In short, animal and in vitro experiments suggest some evidence for photochemical retinal damage upon acute blue light exposure. In humans, a direct evidence of acute light-induced damages to the retina from accidental high-intensity artificial or sunlight exposure is apparent. Ultimately, there is no consistent evidence that long-term exposure to blue light at lower intensity causes any damage to the retina [55]. For even shorter wavelength, ultraviolet radiation, the evidence is more clear [56].

### 4 | THE TIMING OF BLUE LIGHT AND CIRCADIAN DISRUPTION

Circadian rhythms, including melatonin rhythms, are involved in different aspects of facilitation of sleep [57, 58]. Chronic light exposure at the wrong time, at night during shift work for example, may contribute to shifts of the circadian clock phase [59], dependent on duration, wavelength, and intensity of light [60]. While suppressing melatonin secretion during the day is believed to be beneficial, it is not so at night. Circadian disruption is mainly characterized by misalignment between the internal circadian rhythms and the external environmental conditions. Circadian disruption also includes asynchronous SCN with the sleep cycle and with peripheral oscillators in tissues throughout the body, since these peripheral clocks adapt unevenly depending on the tissue [16]. So far, these peripheral circadian clocks, and in some instances their intrinsic zeitgebers, could be identified in almost all investigated mammalian cells [61]. A circadian desynchronization will persist for a variable period of time depending on the exposure pattern and the characteristics of the individual [62], including age and chronotype: morning or evening birds. Light exposure may induce phase advances or delays when applied at different times in the circadian cycle. For instance in humans, light between about 5 AM to 5 PM advances, and light outside this interval delays the circadian clock [63]. Therefore, chronic light exposures at the wrong time may induce phase shift of the circadian system, not allowing for synchronization to the external environmental conditions, and leading to circadian disruption, thus altering both cellular and organ function [61].

When the effects between exposures during the day to blue-enriched white light and white light were compared, the blue-enriched higher color temperature lamps significantly influenced sleep onset (earlier) and reduced sleep latency in the personnel at a research station in Antarctica [64]. This result, suggesting that blue-enriched white light synchronized the circadian rhythm, is corroborated with other results showing that blue light is more efficient in melatonin suppression than other wavelengths [65, 66], and can thus be seen as the most potent zeitgeber. Altered slow wave activity (sleep depth) at the end of the subsequent night of sleep indicated that circadian phase delays might be induced by light exposure before bedtime [43]. These delays are persisting into sleep [43] with a greater effect of blue than green light, suggesting that blue light exposure before bed time can affect sleep. A comparison between the effects of living room light (less than 200 lux) and dim light (about 3 lx) before bedtime showed that even low levels of light in house settings may be sufficient for circadian disruptions in humans: Exposure to room light suppressed melatonin levels and shortened the duration of melatonin production in healthy young subjects [65]. Dim (32 lx), blue depleted LED light, 4 to 7 hours after the subject’s respective waketime appeared to impact the post-illumination pupil response significantly less than regular white LED light, without affecting the melatonin level or subjective sleepiness [67]. Evening exposure to a LED screen, with more than twice the level of blue light (460 nm) emission of a control screen, significantly lowered evening melatonin levels and suppressed sleepiness [68].

For low light levels (40 lx), the melatonin suppression is significantly greater [69] after 2 hours of exposure in the evening to blue-rich (6500 K, CFLs light) than to incandescent light (3000 K). A recent study by Cajochen et al [42] moreover reported, that the alerting effect of light sources with a prominent blue peak appeared to be stronger in the evening and night, compared to morning hours. This alerting effect-exposure timing dependence has further been reported in the context of absent alerting responses during daytime light exposures, suggesting a relationship to circadian features, such as the melatonin level [70]. In another experiment [71] exposing healthy young participants in their natural home environment to
30 minutes of 500 lx blue light an hour before bedtime delayed the onset of rapid eye movement (REM) sleep by 30 minutes. Melanopsin RGCs are extremely sensitive to blue light (see Figure 5) and even exposure to light levels as low as the one from a smart-phone or light emitting e-readers are associated with disruptions of circadian rhythm. Presence of smart-phones in modern life is ever increasing and longer average screen times have been shown to result in shorter sleep duration and worse sleep efficiency [72]. Reading a light emitting e-book before sleep, as compared with printed book, increased the time to fall asleep [73] in young adults (25 ± 3 years old). In individuals reading e-books before bed-time, their circadian clock was delayed, assessed by delayed and reduced phase of rapid eye movement sleep. Melatonin blood concentration levels were suppressed and alertness on the next morning was reduced. Use of the light emitting e-books immediately before bed time also increased alertness at that time. On the other side, daytime blue light exposure has an acute preventive impact on nocturnal light-induced melatonin suppression [74]. Similar tendencies, suggesting increased melatonin suppression and delayed circadian rhythm timing, decreased evening sleepiness and increased morning sleepiness, due to unrestricted light emitting device usage in the evening, have been reported recently [75]. The preventive effect of blue daytime light has proven more efficient than other wavelength composition of daylight. The implications of these findings [65, 68, 69, 72, 73] are, that the melanopsin receptors are particularly sensitive during the evening and nighttime hours, and suggest that many of the sleep disorders may be related to low levels of blue light exposure in the evening, just before bed time. Beside the influence of irradiance it was also shown that color signals, particularly the blue-yellow color discrimination, entrain the inner clock [76]. Narrow band blue light is used today to treat problems such as sleep disorders, jet-lag [77], seasonal affective disorder [78], and premenstrual syndrome. For this, ipRGC photoreceptors are stimulated via light emitting goggles, panels, and other devices.

5 | CIRCADIAN DISRUPTION AND ILLNESS

Chronic disruptions of circadian rhythm may have the potential to seriously affect human health. For instance, decrease of melatonin levels plays an important role in development of chronic diseases and conditions such as cancer [61, 80–83], cardiovascular diseases [83], reproduction [84], endometriosis [84], gastrointestinal and digestive problems [85], diabetes [86, 87], obesity [88], depression [89], sleep deprivation [90], bipolar spectrum disorders [91], and cognitive impairment [92]. However, it is extremely difficult to directly expose the effects of wrong timed light exposures and their long term health consequences, since in modern society expose to artificial lighting is virtually unavoidable. Therefore, epidemiologic studies that suggest light involvement in health risk [93, 94] mainly provide indirect evidence. In general, it is suggested that circadian disruptions are increasing health risk in night-shift workers and flight attendants potentially suffering from both jet-lag and night-shift work. While presence of ambient light during the night sleep is related with circadian disruptions, a decreased health risk is related with long sleep duration and in blind women [95]. Further, results from prospective cohort studies consistently suggest that women with the lowest concentration of the main melatonin metabolite sulfatoxymelatonin, have the highest risk for breast cancer [96, 97]. A recent study could link night shift work with a loss in temporal coordination between the human circadian transcriptome and its environment, most notably by affecting the natural killer cell-mediated immune response and Jun/AP1 and STAT pathways [98].

The most convincing evidence of an association between circadian disruption and health risk is found for breast cancer in night-shift workers. Long periods of night-shift work, which may occur for several years, affects about 10% to 20% of the EU workforce. Prolonged periods of work during the night are the most extreme source of wrong timed exposure to light, which leads to a simultaneous reduction of melatonin production, sleep deprivation and circadian disruption [99]. A recent meta-analysis [100], adjusted for potential confounders in published studies of shift work and female breast cancer risk, demonstrated a significantly increased risk of 40%. Further evidence for the association between shift work and breast cancer [101–105] point toward exposure to light at night as causal factor, but see [106]. A significant association between breast cancer risk and exposure to nonoccupational light during the night at home is found for
women who did not sleep during the period of melatonin level peaks, or who frequently turned on the light during the night [107–109]. Similarly, increased breast cancer risk is also correlated with increasing bedroom light levels. These results are based on self-reports of light exposure and therefore prone to recall bias, which may limit interpretations. Based on evidence in animal experiments on the carcinogenicity of light during biological night, and evidence from epidemiologic studies in humans on the carcinogenicity of light exposure during night work, shift work that involves circadian disruption has been recognized by the International Agency for Research on Cancer (IARC) as probably carcinogenic to humans, Group 2A [82, 110].

It is further hypothesized that the SCN needs repeated input by external, as well as metabolic factors to sustain synchronization between endogenous physiological rhythms and external demands. Modern, hectic lifestyles with omnipresent and energy efficient lightning pollution often lead to a desensitization of the biological clock which results in abnormal endocrine responses, a basic component of type 2 diabetes [87].

Investigations into the cardiovascular system revealed diurnal variations in gene expression, protein expression, and organ function [111] and a potential influence of both internal and intra-organ circadian clock disruptions on the pathogenesis of the cardiometabolic syndrome. A desynchronization of peripheral clocks in cardiac tissue effect the central metabolic activity surrounding cardiomyocytes, thereby leading to accumulated detrimental intracellular long-chain fatty acid (LCFA) derivatives resulting in contractile dysfunctions of the heart [61]. Linking the occurrence of LCFA derivatives with circadian disruption in noncardiac tissue would contribute to several characteristics of the cardiometabolic syndrome [61].

Evidence is also rising that circadian rhythm and both, the irradiance dosis and the spectral irradiance of light, impact eye growth, or emmetropization. Recent reviews shed light on this complex interaction and the interdependence of the disruption of circadian rhythm on the onset and progression of myopia [112, 113]. Clock gene knock-out animal studies are expected to establish further potential associations between circadian disruptions and human pathologies, which are yet to be covered by epidemiological studies [114].

6 | COUNTERMEASURES FOR THE ALTERING EFFECTS OF THE BLUE LIGHT

Blue light is currently considered to have the strongest effect in synchronizing human circadian rhythm. Exposure to low levels of blue of light as well as bright light during night or before bed time may disrupt the circadian rhythm with severe general health implications. At the same time blue light exposure during daytime is crucial for the vitality of the organisms. Retinal illumination decreases with age, for example, due to pupillary miosis, which decrease light transmission, especially in the blue part of the light spectrum. Therefore, inadequate low lighting, especially lacking the blue part of the spectrum, may cause circadian rhythm disruptions. This imposes the importance of proper, bright artificial lighting with a more blue-weighted spectrum during daytime. For example, increasing the blue portion of artificial light may improve performance and learning ability in school kids and employees working indoors, and health will be improved in patients staying at nursing homes or hospitals.

Following the recently accumulated knowledge about circadian rhythm regulation and disruption, it is apparent that there are two key components to keep a healthy circadian system: An increase in the blue portion of the artificial light during daytime should be accompanied with a reduction of the same blue portion of artificial light during the night and evening hours. Electronic manufacturers and software providers already offer a variation of blue light blocking features for displays. While the implementations in general decrease the amount of blue light emitted by the devices, the remaining share of short wavelengths in white light is not accounted for. Importantly, it has been reported that a pure blue light filter has an insufficient effect on melatonin suppression without an accompanied brightness reduction [115]. A simple solution that may effectively block and reduce the blue portion of the light spectrum before bedtime are blue light blocking glasses and lenses [116, 117]. Wearing blue light blocking eyewear before [118, 119] and during [120] bed time may effectively attenuate LED induced melatonin suppression [121, 122] and thus can potentially facilitate the adaption to new social schedules [123] and reduce sleep disturbances and their consequences among the general population. Further, a reduction of the blue portion in artificial light during nighttime hours could protect shift workers against disorders such as cancer and cardiovascular disorders. Importantly, all solutions should consider the optimal spectral requirements of both, conscious and unconscious photo-reception.

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CONFLICT OF INTEREST

S.W., M.E., and I.V.I. are researchers at the University of Tuebingen; S.W., P.S., C.L., and I.V.I. are employed by Carl
Zeiss Vision International GmbH, manufacturing spectacle lenses and sunglasses. There is no conflict of interest regarding this study.

**AUTHOR CONTRIBUTIONS**

S.W. and I.V.I. did the literature study and analysis; S.W., M.E., and I.V.I. wrote the manuscript. P.S. and C.L. designed the illustrations; all reviewed the illustrations and all reviewed critically the manuscript.

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