Slowly seeing the light: an integrative review on ecological light pollution as a potential threat for mollusks

Ahmed A. A. Hussein 1,2,3 • Erik Bloem 3 • István Fodor 4 • El-Sayed Baz 1 • Menerva M. Tadros 2 • Maha F. M. Soliman 1 • Nahla S. El-Shenawy 1 • Joris M. Koene 3

Abstract
Seasonal changes in the natural light condition play a pivotal role in the regulation of many biological processes in organisms. Disruption of this natural condition via the growing loss of darkness as a result of anthropogenic light pollution has been linked to species-wide shifts in behavioral and physiological traits. This review starts with a brief overview of the definition of light pollution and the most recent insights into the perception of light. We then go on to review the evidence for some adverse effects of ecological light pollution on different groups of animals and will focus on mollusks. Taken together, the available evidence suggests a critical role for light pollution as a recent, growing threat to the regulation of various biological processes in these animals, with the potential to disrupt ecosystem stability. The latter indicates that ecological light pollution is an environmental threat that needs to be taken seriously and requires further research attention.

Keywords
Artificial light • Biorhythm • Mollusca • Reproduction • Snails • Slugs • Zeitgeber

Introduction
Natural light is known to be a crucial regulating cue for the biological world and generally acts as a zeitgeber for biological rhythms (Bradshaw and Holzapfel 2010; Foster and Roenneberg 2008; Ragni and Ribera D’Alcalà 2004). As a natural abiotic factor, it is known to influence many behavioral and physiological processes in animals, e.g., reproduction, energy storage, and (neuronal) activity. One important aspect is the seasonal change in light conditions, meaning that even though natural light is not constant but varies over time, this still provides sufficient information to entrain biological rhythms.

Short-term variation in natural light can, for instance, be due to the presence of clouds that block part of the light coming from the sun or stars, and light intensity may change rapidly with increasing sky turbidity (Cronin et al. 2014). As a result, the natural light intensity of the sun can range from 120,000 lx for direct sunlight at noon to less than 5 lx during misty sunsets or sunrises (Gorman et al. 2005). Despite this variation, such light information still serves as a zeitgeber for many of the behaviors that depend on a circadian or circannual rhythm. Nevertheless, in recent years, it has become clearer that the use of artificial light, as part of increased human activity in environments, can affect or even shift the natural rhythmicity of animals (Gaynor et al. 2018).

Nevertheless, in contrast to urbanization effects caused by chemical pollution (Likens et al. 1996), habitat restructuring (Poff et al. 1997), and invasive species (Ricciardi and Rasmussen 1998), those effects caused by light pollution have only been recognized in the past years (Longcore and Rich 2004; Moore et al. 2006; Nightingale et al. 2006). Insects and larger (vertebrate) animals have received attention on how they are affected by such light pollution. However, the second largest group of animals, the mollusks, have been largely

1 Zoology Department, Faculty of Science, Suez Canal University, Ismailia 41522, Egypt
2 Theodor Bilharz Research Institute (TBRI), Giza, Egypt
3 Department of Ecological Science, Faculty of Science, Vrije University, De Boelelaan 1085, 1081 Amsterdam, Netherlands
4 NAP Adaptive Neuroethology, Department of Experimental Zoology, Balaton Limnological Institute, Centre for Ecological Research, 8237 Tihany, Hungary
Artificial light and light pollution

The quantity of artificial light can be used as a rough indication for the size and development of contemporary human societies (Cinzano et al. 2001) because the use of light at night has turned out to be fundamental for modern societies (Höcker et al. 2010a). When thinking about such artificial light at night (often abbreviated as ALAN), one should not only think of city lights at night but also lights from traffic, greenhouses, and agricultural systems, for example. In addition, due to the development of the world economy, industrial facilities such as ports, railway yards, and airports are illuminated all the time, as are lit marketing and advertising columns. All these examples of sources of artificial light could increase environmental stress and alter the natural light-dark cycle of organisms (Barré et al. 2020; Baz et al. 2013; Bruce-White and Shardlow 2011; Dominoni et al. 2020; Health Council of the Netherlands 2000; Höcker et al. 2010a, 2010b; Longcore and Rich 2006, 2004; Navara and Nelson 2007; Outen 1998; Perkin et al. 2011; Verheijen 1985, Verheijen 1960).

The most common definition used for light pollution is the change of natural light patterns in the night environment caused by the introduction of artificial light. Höcker et al. (2010b) showed that the use of artificial lighting is spreading at 6% every year. This can be in the form of direct illumination of the environment surrounding the light sources but can also happen through sky glow resulting from this illumination. Such artificial sky glow can expand the ecological impact of light pollution, as a side effect, to many miles beyond cities. The impressive increase in the use of artificial light at night has put light pollution on the list of threats to ecosystems. Many studies confirm that lighting at night affects wildlife including plants, invertebrates, fish, amphibians, reptiles, birds, and mammals (Barré et al. 2020; Davies et al. 2014; Dominoni et al. 2020; Gaston et al. 2013; Gaston and Bennie 2014; Höcker et al. 2010b; Longcore and Rich 2006; Longcore and Rich 2004), and may have consequences for biological processes (Lewanzik and Voigt 2014). For example, a recent study investigating the mechanisms underlying the near-perfect synchronization of fireflies’ glow also pointed out that some of these species, which use their glow to attract mates, have found themselves competing for attention with human sources of light (Sarfati et al. 2020). Our conceptual framework (Fig. 1) shows how light pollution can affect these processes in terms of changing natural light photoperiod, wavelength, and intensity.

Quantifying light pollution

One important aspect of light pollution is its quantification. First of all, it is important to differentiate between astronomical light pollution and ecological light pollution, as pointed out by Longcore and Rich (2004). They define astronomical light pollution as the reduction in the visibility of stars originating from artificial light sources at night, a phenomenon also known as sky glow. Next to this, they referred to the effects caused by changes in the natural light levels at night as “ecological light pollution.” In the following, we consider ecological light pollution only.

The quantification of natural light frequently includes measuring the brightening at a given place. There are many ways to express such brightening, but the most widely recognized is the measure of light occurrence per unit of area. Measuring light usually depends on two properties of light: intensity and spectral composition. Nowadays, the most commonly used unit to measure light is lux, which takes the intensity of light into account but neglects the type of light (i.e., the spectral composition represented by wavelengths). Longcore and Rich (2004) suggested that light should be quantified by counting all properties of light. Scientists are therefore advised to quantify brightening in photons per square meter every second accompanied by the wavelengths of that light. This clearly

overlooked so far. Nevertheless, such animals do use light as a zeitgeber as well. For example, in the freshwater pond snail, Lymnaea stagnalis, circannual changes in environmental light conditions can affect reproduction, energy storage, and neuronal activity. For this species, extended photoperiods are linked to increased glycogen storage and the initiation of overwintering dormancy (Bailey 1981; Hemminga et al. 1985). Such seasonal changes have also been observed in the synaptic connections between the well-studied RPeD1 neuron of L. stagnalis and its follower cells (Copping et al. 1999), as well as in how well these snails can deal with anoxic conditions (Buck et al. 2017). One of the very few studies addressing the effects of light pollution in mollusks concluded that their behavioral changes could potentially disrupt interspecific interactions, and thereby ecosystem functioning (Underwood et al. 2017).

In this review, we aim to specifically focus on the potential effects of light pollution on mollusks to inspire and guide research in this direction. To do so, we first define what the term “light pollution” means exactly. Then, we provide a brief overview of the different roles that natural light can play, using some known examples from the animal kingdom and asking whether there is any evidence for this in mollusks. In each section, we also specifically focus on what is known about light and its perception in mollusks (summarized in Table 1) and in which ways these animals can (potentially) be affected by light pollution. This has led us to a conclusion in which we highlight what we think are the most fruitful areas of research to answer some of the pertinent questions.

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| Species                  | Light condition          | Behavior or process | Interaction | Effect                  | Reference                                |
|-------------------------|--------------------------|---------------------|-------------|-------------------------|------------------------------------------|
| *Aplysia californica*   | Short photoperiods       | Egg laying          | Temperature | Increase                | (Wayne and Block 1992)                   |
| *Helix aspersa*         | Light                    | Juveniles growth    | Temperature | No effect               | (Benbellil-Tafoughalt et al. 2009)       |
| *Cornu aspersum (Helix aspersa)* | Choice between light & dim light | Attraction | Prefer light over dim light |                          | (Perea et al. 2007)                      |
| *Helix aspersa var. maxima* | Long photoperiods       | Egg laying          | Temperature | Increase                | (Gomot 1990)                              |
| *Limax maximus*         | Absence of light         | Weight (growth)     | Temperature | Increased               | (Jess and Marks 1998)                     |
| *Limax valentianus*     | Long photoperiods        | Growth rate         | Temperature | Higher                  | (Sokolove and McCrone 1978)              |
|                         | Long photoperiods        | Egg production      | Temperature | Start sooner            | (Hommay et al. 2001; Sokolove and McCrone 1978; Udaka et al. 2008) |
|                         | Short photoperiods       | Egg size            | Temperature | Larger                  |                                          |
|                         |                          | Gonads              |             | Heavier and quicker development |                                          |
| *Lymnaea acuminate*     | Short photoperiods       | Growth rate         |             | Higher                  |                                          |
| *Lymnaea stagnalis*     | Red light                | Fecundity, hatchability, and survivability | Fed with chlorophyll | Reduced | (Kumar et al. 2016) |
| *Physa integra*         | Light                    | Attraction           |             | Yes                     | (Clampitt 1974)                          |
| *Physa pomiliia*        | High intensity           | Attraction           |             | Yes                     | (Badman 1966)                            |
limits the usefulness of lux, especially if one considers that organisms can detect and recognize light at different wavelengths than those visible to humans. Therefore, light with the same intensity but with different wavelengths may have a very different impact on an organism because it depends on that species’ spectral detection range (Dominoni et al. 2020). For example, moths are attracted to high-pressure sodium lights but not to low-pressure sodium lights; these lights have the same lux value but only high-pressure sodium lights produce the ultraviolet wavelength that attracts moths (Rydell 1992).

**Light perception**

The previous section illustrated that the physical properties of light can affect organisms in different ways. Therefore, it is important to get a good sense of how organisms use light and darkness as a resource (Gerrish et al. 2009; Kronfeld-Schor and Dayan 2003). The direction, period of exposure, and physical characteristics of light provide organisms with vital information about their surrounding environment, such as locations, the timing of days, different seasons, and years (Neff et al. 2000; Ragni and Ribera D’Alcalà 2004). The usefulness of this information also depends on how efficient organisms are at detecting and recognizing light. Different visual mechanisms have evolved and the level of specialization of visual organs and the accompanying photoreceptors can range from receptors located on the body wall to well-developed eyes (see details in Gehring and Seimiya 2010; Gehring 2014). The origin of vision is assumed to be found in the terrestrial cyanobacterium, *Leptolyngbya* sp., in the form of an eyespot or “stigma” that consists of long, slender trichomes (filaments). This eyespot allowed these bacteria to be positively phototactic, thus enabling them to move toward a light source. Besides, the eyespot is characterized by containing carotenoid-rich lipid globules that are also found in phototoxic flagellated algae (Albertano et al. 2000).

In general, one of the main reasons for the efficiency or deficiency of vision is related to animal photoreceptor cells and their associated pigments. One important pigment involved in vision is opsin that mediates the conversion of a photon of light into an electrochemical signal as the first step in the visual transduction cascade. The photoreceptor cells in animal eyes are often classified into microvillar cells with rhodomeric opsin (r-opsin) and ciliary cells with ciliary opsin (c-opsin) according to whether the sensory surface is enlarged by microvilli or by cilia, each type having specialized molecular characteristics (Döring et al. 2020). In response to light, rhodomeric photoreceptor cells in protostome eyes are known to signal via the G\(_{\alpha q}\)-mediated inositol 1,4,5-triphosphate (IP3) cascade opening transient receptor potential (TRP) ion channels in the photoreceptor cell membrane that leads to a depolarization (Fain et al. 2010). In contrast, ciliary photoreceptor cells of vertebrate eyes are known to signal via the G\(_{\alpha i/t}\)-mediated cGMP cascade closing cyclic nucleotide-gated (CNG) channels and leading to a hyperpolarization (Nilsson and Arendt 2008; Wensel 2008).

The presence of these two types of receptor cells has been suggested to already be present in the eyes of the bilaterian ancestor for the following main reasons. First, both photoreceptor cells are found in protostome and deuterostome animals and they have distinct molecular characteristics (Arendt 2008; Arendt et al. 2004; Arendt et al. 2002; del Pilar Gomez et al. 2009; Gehring 2014; Panda et al. 2002). Second, c-opsin in protostomes seems to be present only in extraocular photoreceptor cells of certain groups of annelids (Arendt et al. 2004) and arthropods (Beckmann et al. 2015; Velarde et al. 2005) and to be lost in all other protostomes (Döring et al. 2020; Ramirez et al. 2016; Vöcking et al. 2017). This supports the assumption of ancestral extraocular expression of vertebrate c-opsins in brain extraocular photoreceptors (Arendt et al. 2008; Arendt et al. 2004; Shubin et al. 2009) and suggests that the involvement of c-opsins in the visual cells of cerebral eyes evolved later, most likely a chordate-specific phenomenon (Vopalensky et al. 2012).
However, the recent discovery and investigation of a new type of visual opsins, xenopsin, have pointed toward a more complex situation. Xenopsin is present in ciliary photoreceptor cells of a wide range of protostome taxa (summarized in Döring et al. 2020) and although it shares important functional sequence motifs with c-opsins, they do not group in phylogenetic analyses (Ramírez et al. 2016; Rawlinson et al. 2019; Vöcking et al. 2017). Until now, the notion was that xenopsin and c-opsin are mutually exclusive in a given species; however, a new study refuted this view reporting the first organism, the annelid Malacoceros fuliginosus, that has both xenopsin and c-opsin (Döring et al. 2020). Furthermore, photoreceptor cells (being potentially polymodal sensory cells) expressing both xenopsin and t-opsin and exhibiting both microvilli and cilia have been found in larva of the mollusk Leptochiton asellus and in larva of the annelid Malacoceros fuliginosus (Döring et al. 2020; Vöcking et al. 2017).

In the light of these new findings, Döring et al. (2020) have provided a new perspective for comparative eye research: for example, highlighting that xenopsin is an important visual pigment in protostomes and that ciliary eye photoreceptor cells may not be of the same evolutionary origin in protostomes and deuterostomes. As a result, they proposed all conceivable alternative scenarios for the evolution of these opsins and photoreceptor cells in bilaterian animals, which clearly indicated that the exact evolutionary processes remain to be determined and further studies are required for a better understanding.

We will here review some details of the light perception and vision of the Gastropoda (within the phylum Mollusca, the second largest animal group in terms of species number, after insects). The Gastropoda is an extremely diversified class, and its number of species is estimated to lie between 65,000 and 80,000 living snails and slugs. They are often used as bioindicators for the quality of the environment and to detect the effects of different types of pollutants on ecosystems (Bouchet et al. 2005). These species can vary greatly in their behavior, reproduction, habitat, anatomy, and mode of obtaining food. The eyes of a gastropod can be situated at the base of the tentacles, on short stalks, or at the end of their retractable tentacles, respectively, Basommatophora (Hygrophila), and Stylommatophora.

Besides ocular receptors in the eyes, gastropods also have non-ocular light receptors and they use both to avoid total darkness and high intensities of illumination (Brown and Brown 1973; Gotow and Nishi 2009; Lyons et al. 2006; Rossetti and Cabanac 2006). The eye is used for phototaxis and for regulating the behavioral patterns on a daily and seasonal basis. It is still unknown to what extent gastropods can use their eyes to distinguish different shapes and forms. However, they have been shown to distinguish between checkerboard patterns in black and white, as well as a gray background, and they can distinguish between horizontal and vertical lines (Chase 2002). This pattern detection can be explained by the presence of different classes of neurons. For example, Stoll and Bijlsma (1973) presumed that there were 2 classes of neurons in the eye of L. stagnalis: photoreceptors and optic ganglion cells. A later morphological examination suggested that there might be 3 types: photoreceptors I and II, and optic ganglion cells (Bobkova 1998). This distinction is not based on the size and area of each type of photoreceptor in the retina but rather on microvilli. The photoreceptor axons extend out of the eye to the central nervous system where they connect with the terminal branches of the statocyst hair cells at a synaptic contact (Sakakibara et al. 2005). It is unclear whether these statocyst hair cells, which are generally involved in balance and orientation (Janse et al. 1988), are also sensitive to light themselves or whether light and other visual information are integrated at this point.

The non-ocular light receptors are known to mediate behavioral responses like the shadow reflex, the defense response when there is a sudden drop in illumination such as that caused by a predator (Ramírez et al. 2011). These non-ocular receptors are distributed over the body wall. Moreover, there is some evidence that when the skin is thin enough, and the light intensity high enough, central neurons can directly be triggered by light (Chase 2002). This direct triggering is hypothesized to be mediated by light-sensitive carotenoids and/or other photopigments that may be present in intracellular neuronal organelles (Gotow and Nishi 2009; Ter Maat et al. 2012) but remains to be demonstrated.

Previous investigations have demonstrated that L. stagnalis has TRP-channel-mediated ocular photoreceptor cells and CNG channel-mediated non-ocular photoreceptors. The latter is found especially around the mantle and foot. However, CNG and TRP photoreception can also act synergistically. This happens in the marine gastropod Onchidium verruculatum that has additional visual and dermal photoreceptors situated in the dorsal eye and eyestalk. Furthermore, L. stagnalis has been confirmed to have three dermal photoreceptors, one in which cyclic guanosine monophosphate acts as a second messenger in the dermal photoreceptor, a second type that contains rhodopsin as a photopigment, and a third that uses the photosensitive protein, Arrestin (Takigami et al. 2014).

The roles and effects of natural and artificial light on organisms

Organisms use light in many different ways, for example as a resource and also for regulating activity patterns like sleep, reproduction, growth, and orientation. Depending on the kind of physiological process or behavior activity, either light or its absence is a prerequisite. The availability of periods of exposure to light and darkness determines the time that is available
for each process. As a result, a disruption in the natural availability of light and darkness can have positive or negative consequences for organisms depending on whether either is a limiting factor.

### Light as a resource and activity determinant

The use of light as a resource is found in the sea slug *Elysia chlorotica* whose metabolism relies on photosynthesis by an alga. This slug feeds on the intertidal alga *Vaucheria litorea*. Instead of digesting the entire algal cell, it conserves the chloroplasts in its gut where they fuse with gut cells and continue photosynthesizing (Mujer et al. 1996). Interestingly, some *Elysia chlorotica* slugs have even reported having the capacity to utilize photosynthesis for up to a year after just a couple of feedings. Such use of light as a resource may provide an example where light pollution may be beneficial since more exposure to light would provide the slug with more energy through photosynthesis (Rumpho et al. 2008).

Light also affects activity patterns of many organisms, with some being active during the day while others being nocturnal (Kronfeld-Schor and Dayan 2003), a separation of activity between organisms that is at least in part due to survival (Gutman and Dayan 2005). However, biological and developmental investigations have concentrated on diurnal organisms, a significant proportion of species adjusted to be active at night during low-light or dark conditions (Hölker et al. 2010b; Lewis and Taylor 1965). These nocturnal organisms seem to be more affected by changes in natural light levels at night in terms of activity, behavior, and survival. Because artificial lighting generally reduces darkness to a semi-darkness level that may be similar to sunset or moonlight conditions (16 L:8D) produce 2- to 3-fold more eggs than normal-day snails (12 L:12D). To confirm their findings on the

### Habitat choice and biodiversity

Organisms can adapt to changes in natural light levels and durations in their environment by leaving light-polluted areas to occupy new habitats. This may make the species invasive in such a new habitat and/or may affect population density levels of their own or native species. González et al. (2014) reported that the occupation level of communities of sandy beach beetles increases with the quality of the sky at night and is thus negatively affected by light pollution due to urbanization. These findings are partly in agreement with those for the impact of light pollution on the amphipod *Orchestoidea tuberculata* (Fanini et al. 2016; Giacconi 2006; González et al. 2014). For black-tailed godwits, there is also suggestive evidence that the nest location depends on the amount of surrounding light, with preferred nesting sites far away from road lighting (De Molenaar et al. 2000). Also, investigations on bats in Sweden demonstrated that artificial night lighting is the reason for a change in bat biodiversity; however, common species remained, but rarer species diminished even more in abundance (Rydell 1992). Finally, within the Mollusca, there is also evidence that the presence of light influences habitat choice. Perea et al. (2007) reported, in a study on the species *Cornu aspersum* (formerly *Helix aspersa*), that these snails preferred light over dim light. The latter finding is in agreement with an earlier study by Badman (1966) in which he had found that another species, the freshwater snail *Physa pomillia*, was attracted to the high intensity of light.

### Reproduction

Reproduction is another essential process that can be impacted by changes in natural light patterns in various ways. Females of the frog species *Physalaemus pustulosus* choose a mate more quickly when exposed to higher levels of light, probably to avoid the risk of predation (Rand et al. 1997). In addition to mate choice, the frequency of breeding can also be affected—as Longcore and Rich (2004) found out when studying the effect of sky glow resulting from stadium lighting during football games. In their semi-field experiment, frogs were found not to mate during this lighting period, but they resumed mating after the lighting was blocked by covering their enclosure. Furthermore, Mc Lay et al. (2018) found that *Drosophilia melanogaster* flies exposed to 10 lx before mating courted longer than flies exposed to darkness at night before mating. They also found that female oviposition patterns differed between the two light treatments and explained this by determining that female flies exposed to dim light at night had a lower level of reactive oxygen species (ROS) in their ovaries than those exposed to 0 lx.

Egg laying behavior is one of the reproductive parameters that is often influenced by light periods and light pollution. For example, Ter Maat et al. (2007) reported that the period of exposure to light has an effect on egg laying in the snail *L. stagnalis*. They found that snails exposed to long-day conditions (16 L:8D) produce 2- to 3-fold more eggs than normal-day snails (12 L:12D). To confirm their findings on the
influence of light on egg laying behavior, they followed up this study by adding the so-called clean water stimulus as an additional factor, which has been shown to induce egg laying when snails are transferred from dirty to clean water and/or jar. They found that when this stimulus was given in the dark, egg laying was induced significantly less than in the light. Hence, light seems to help to stimulate the caudal-dorsal cells (CDCs) that are responsible for releasing the egg laying hormone (CDCH). Interestingly, the snails did not need their eyes to exhibit this difference, so while it remains elusive how light reaches the CDCs it seems likely that this happens via non-ocular photoreception (Ter Maat et al. 2012).

In contrast to freshwater snails, Wayne and Block (1992) stated that for the marine slug *Aplysia californica* exposure to different photoperiods only has a minor influence on reproduction, the main controlling factor being temperature. They found that animals that were obtained in autumn and kept in warm water laid eggs more frequently than those in cold water, regardless of photoperiod. This is directly opposed to what Gomot et al. (1989) found in their study on the terrestrial snail *C. aspersum*, which may be down to differences in habitat (i.e., marine vs. terrestrial). The latter study concluded that light is a dominant signaler for inducing egg laying behavior, based on their finding that egg laying is influenced by both temperature and light, but that long-day snails produced more eggs than short-day ones. Moreover, egg laying stopped after 6 weeks of exposure to 15 °C at short days while it continued under normal conditions with 15 °C and long days (which is in agreement with Stephens and Stephens (1966)). Therefore, they conclude that egg-laying hormone production is stimulated by light as in *L. stagnalis*.

Besides egg laying itself, other reproductive parameters can also be affected by light. Kumar et al. (2016) found that the fecundity, hatchability, and survivability of *Lymnaea acuminata* were reduced after being fed with chlorophyllin and exposed to red light. Hommay et al. (2001) also reported that egg production started sooner, more eggs were laid, that these eggs were larger, and that their hatching was significantly higher under long photoperiod when compared to a short photoperiod treatment.

One more reproductive parameter that is greatly affected by light is the growth and development of gonads. Sokolove and McCrone (1978) previously found that *Limax valentianus* slugs that were held under a short photoperiod had gonads and oocytes that were heavier than those from slugs held under long photoperiod. This difference was developmental because it was apparent 90 days after hatching, but disappeared after 120 days and the short photoperiod slugs also reached the last stage of spermatogenesis (Hommay et al. 2001; Udaka et al. 2008).

**Orientation and communication**

Light is also an essential player in the orientation of animals, for example to hide from enemies or predators, to locate mating partners, to find food, and/or to migrate; any change in the natural light pattern may alter their perception of direction (Baker 1990). The effect of how migration is affected by artificial lighting at night is found in migratory birds. For example, artificial outside lighting was found to disturb the orientation of young birds, especially in cloudy conditions (Abt and Schultz 1995). Moreover, some migrating birds have been found to fly near lights under bad weather conditions and may get disoriented or even trapped in light areas (Evans Ogden 1996).

Both diurnal and nocturnal animals are affected by artificial light in their habitat. The increased illumination at night may enhance the ability of diurnal animals to orient themselves and may alter certain behaviors such as foraging in birds (Hill 1992) and reptiles (Schwartz and Henderson 1991). Negative effects of artificial light at night are especially experienced by nocturnal animals because they are adapted to navigate better under dark conditions (Park 1940). One famous example comes from the hatchlings of sea turtles that move from their nests on sandy beaches toward the ocean guided by the natural light, but with the increase of artificial lighting surrounding beaches, the hatchlings get disoriented, and may even move opposite to the direction of the shoreline (Salmon et al. 1995).

By using illumination, animals may also be attracted to or repelled by light sources (Health Council of the Netherlands 2000). A lot of species of insects are attracted to illumination, such as moths. Other taxa like lacewings, beetles, bugs, caddis flies, crane flies, midges, hoverflies, wasps, and bush crickets exhibit similar attraction to light (Eisenbeis and Hassel 2000; Frank 1988; Kolligs 2000). This behavior increases the risk of being predated by bats and spiders (Kiefer et al. 1995; Rydell 1992). In contrast, some nocturnal spiders are negatively phototaxic and repulsed by light (Nakamura and Yamashita 1997) while other insects are positively phototaxic (Summers 1997). In other species the attraction or repulsion effect of light may be used in more applied ways, such as using lights to attract fish to ladders—artificial passages that allow them to bypass dams and power plants (Haymes et al. 1984).

For Mollusca, there is some evidence that light attracts snails (van Duivenboden 1982) and plays a role in predator avoidance (Pankey et al. 2010). *L. stagnalis* can escape from predators via the well-known whole-body withdrawal response. According to behavioral and physiological research, exposure to shadow stimulates slams and their predators in the opposite way. Shadow stimulates the predator to attack, whereas it stimulates non-ocular photoreceptors in the slams to send alert signals to the left and right pedal dorsal 11 neurons. These neurons connect to motor neurons 13–16 via chemical synapses and can initiate the escape behavior (Takigami et al. 2014). This species is even attracted to light when the eyes are experimentally removed (van Duivenboden 1982). The latter clearly indicates that non-ocular
photoreceptors are involved. Such positive phototaxis is also seen in the freshwater pulmonate Physa integra, which moves toward the shore in spring and is guided by light (Clampitt 1974). Clampitt (1974) found that these snails predominantly moved toward the light and this choice seemed independent of a gravitational cue (Clampitt 1974).

Communication via direct visual cues and/or signals is also used by organisms and may be disturbed by artificial lighting at night. The increased lighting at night in the environment of glowing worms interferes with the attraction of mates via bioluminescence (Lloyd 1994). Hence, the presence of artificial lighting at night decreases the chances of glowing female worms to be located and fertilized by males. So far, there is no evidence from the literature suggesting that light pollution affects sexual signals in mollusks. However, this may not seem surprising for gastropods, which often have limited visual abilities (Di Cristo and Koene 2017). However, it may affect cephalopods since these have well-developed eyesight and are attracted to light. So, this remains a research area that deserves more attention in future studies.

**Disturbance of the circadian clock**

There is a crucial role that light plays in synchronizing the nervous system to the external 24-h day-night rhythm (e.g., Baz et al. 2013). The timing of daily rhythms is regulated by an endogenous timekeeping system referred to as the circadian clock. In mammals, this is located in the suprachiasmatic nucleus and in mollusks in the eyes. The biological rhythms differ between organisms and are influenced by periodic cycles (such as day/night, season, high/low tide, and lunar cycle) as well as temperature, wind, and feeding, etc. Many biological functions are linked to such daily or annual periodicity and are controlled by the neuroendocrine system via the epiphysis. This gland is located in the brain and responsible for stimulating the hormonal pathway that produces melatonin (De Molenaar et al. 1997). Artificial illumination at low intensity during the night has been shown to alter secretion of melatonin and thereby internal physiological functions in many species, such as birds, fish, and mammals (Bedrosian et al. 2011a, 2011b; Cos et al. 2006; Evans et al. 2007; Navara and Nelson 2007).

Many studies were done to evaluate the effect of altering melatonin production due to exposure to light pollution. In humans, some researchers have reported a negative relationship between disturbance of melatonin production and cancer risk for people structurally working during the night (Megdal et al. 2005; Reiter et al. 2011; Stevens 2009). At very low-intensity levels of illumination, melatonin secretion may be affected. Rats exposed to illumination intensity of 0.2 lx during the night had decreased levels of melatonin production (Dauchy et al. 1997), similar to the effect of 1 lx on hamster (Brainard et al. 1982), and a higher rate of tumor growth and immune system inhibition were observed (Bedrosian et al. 2011b; Dauchy et al. 1997).

Desynchronization of the biological clock can be one of the impacts that light pollution has (Health Council of the Netherlands 2000). The disturbance may lead to insufficient rest or sleep which may consequently influence fitness and/or survival. Black-tailed godwits and bearded tits can no longer adjust their digestive system for feeding on seeds during winter in Africa from being adjusted to insect feeding during summer due to a lack of synchronization (De Molenaar et al. 2000).

There is no published evidence on the effect of light pollution on the circadian clock of mollusks and/or the secretion of melatonin. This may be because light pollution has only recently been recognized as an environmental threat. Nevertheless, at least one study hypothesized that there are negative effects of disturbance of circadian rhythms of snails due to sky glow resulting from artificial light (Lyrytimäki et al. 2012).

**Growth**

Growth is also impacted by light pollution. For example, laboratory and field studies by Luarte et al. (2016) revealed that locomotor activity, foraging behavior, and growth rate of the amphipod Orchestoidea tuberculata were highly affected by light pollution. In the field, they showed that low light (60 lx) reduced amphipod feeding and growth rates. These findings are in agreement with work showing that light pollution reduces consumption rates in rodents (Vasquez 1994) and decreases the development of juvenile and suckling bats (Myotis emarginatus and M. oxygnathus) (Boldogh et al. 2007). Similar results are also shown for talitrid amphipods’ growth rate (Duarte et al. 2016).

Raap et al. (2016a, 2016b) tested the effect of artificial light at night on the physiological parameters (body mass and oxidative status) during development, using nestlings of a free-living songbird, the great tit (Parus major). They measured multiple biomarkers after two nights of exposure to 3 lx 2 h before sunset and 1 h after sunrise of the following morning. They found that light inhibits body mass but no difference in the oxidative profiles of the exposed individuals. However, this investigation provides evidence that artificial light at night may negatively influence the growth of free-living nestlings that may persist throughout adulthood.

In mollusks, there is also the potential for light pollution affecting growth and development. Ter Maat et al. (2007) discovered that a relationship exists between the daily duration of exposure to light and the growth speed and amount of energy stored in the freshwater pulmonate L. stagnalis. Furthermore, the amount of stored energy was higher in the medium-day snails than those in the long-day snails. This is as expected because in spring and autumn food availability is lower; therefore, it is advantageous to store energy whereas in summer food availability is high and there is thus no need to...
store energy. With a decreasing amount of food available, the dry-weight density of the long-day snails decreased.

For another mollusk, the land snail *Helix aspersa var. maxima* weight was used as an indicator of growth. The weight increased by 36% in the absence of light at 15 °C compared to snails exposed to 16 h light, while at 20 °C and a light period of 16 h, the weight improved by 11% compared to those reared in total darkness. However, at 20 °C, snails were larger in weight by 91% than those raised at 15 °C independent of their photoperiod (Jess and Marks 1998). This finding does not agree with the findings from Benbellil-Tafoughalt et al. (2009) who reported that the growth of juveniles of *Helix aperta* was influenced only by temperature and that exposure to different photoperiods had no effect.

For the terrestrial slug species investigated, growth was greatly affected by exposure to different photoperiods, but in different directions. In *Limax valentianus*, slugs were heavier and had a higher growth rate under short photoperiod (12 L:12D) than those held under long photoperiod (16 L:8D) (Hommay et al. 2001; Udaka et al. 2008). In contrast, *L. maximus* gained more weight in long photoperiods (16 L:8D) compared to short photoperiods (8 L:16D). These opposing findings clearly highlight that the underlying mechanisms may differ, even between closely related species.

### Conclusions and future perspective

The growing use of artificial light at night, such as street lights, greenhouses, industrial facilities, and advertising columns, has the potential to increase the exposure of both aquatic and terrestrial organisms to continuous 24-h photoperiods. This increase could be accompanied by light intensities and spectral compositions, but the real impact of the biological and ecological consequences of artificial night lighting is still unknown. However, researchers are starting to uncover that outdoor illumination affects biological rhythms, and there is a clear need for further exploration of the impact of light pollution on biological systems. Gastropoda seems a suitable class of animals for studying the possible impacts on ecosystems because many members of this group are impacted by changes in natural light regimes (Table 1). As one of the main mollusk can model species, *Lymnaea stagnalis* seems highly suitable for testing the effects of light on reproduction, growth, survival rate, and development success because of its demonstrated sensitivity to different light conditions. For this species, exposure to longer photoperiods is already known to enhance and/or initiate various biological processes, such as reproduction (Ter Maat et al. 2012; 2007). Importantly, more work needs to be done to establish whether not only extended constant photoperiods but also lower levels of artificial light at night disturb other processes such as movement activity, behavior, feeding, and ability for learning and memory formation. The latter neurobiological processes involve neuropeptides produced by neuroendocrine cells in the relatively simple central nervous systems (CNS) of *Lymnaea stagnalis*. Their simplicity and well-mapped CNS should enable researchers to better understand the mechanisms responsible for the expected impacts of light pollution on different behavioral and biological processes. In addition, the extensive knowledge about the underlying regulatory mechanisms and the availability of genome and transcriptome data for this species will facilitate interpretation (Fodor et al. 2020).

Earlier research has indicated that light can also have effects that interact with other factors, such as temperature and food availability. While in terrestrial gastropods light seems to be the main trigger, such interaction effects are still largely unexplored in aquatic species and deserve attention in the future, especially given that water temperatures are predicted to rise. Also, from prior research, a primary issue emerges: what consequence does light pollution have on gastropod population density? And does this differ when food is abundant and when food is a limiting factor? Ter Maat et al. (2007) showed that the availability of food and the presence of a longer photoperiod together have a positive effect on the development and reproduction of *L. stagnalis*. This would predict a rise in population density, with the potential to trigger a situation where the species becomes a pest if light pollution continues to expand. The gathering of snails around light sources may increase predation risk, just as it does in moths (Frank et al. 2006). Furthermore, the continued exposure to light may condition gastropods to stop their shadow reflex because of the large amount of false triggering. Combining the latter two might then result in decreased population density, but this also remains to be shown and/or experimentally tested. Hence, the need for further investigation of the effects of a 24-h light period on population dynamics of the freshwater snail through reproduction and behavioral responses like the shadow response, movement activity, and learning becomes necessary for better understanding the implications.

We aim to conduct such research in our laboratory in future. Most reviewed studies only decreased or increased the length of the photoperiod, so it is fair to assume that they used an average light intensity that the gastropods experience in their natural habitat or their culturing facility. Hence, the question remains whether light pollution has a strong enough light intensity to cause a similar effect on these snails as an extended photoperiod at normal intensity. This indicates that in certain animals a low intensity of light pollution is sufficient to change their behavior, so it is entirely possible that gastropods are also affected by such levels of sky glow light pollution and thus remains worth testing. Such research should then focus also on quantifying the lowest level of light necessary to evoke a change in behavior, which will also help to establish the safe limit of light exposure at night in terms of intensity, spectrum, and duration.
Finally, further exploration of this topic will increase our empirical knowledge and help in better understanding the possible impacts of light pollution. Identification of the pigments involved in light perception in mollusks (and animals in general) will also contribute to a more complete understanding of the mechanism and molecular networks underlying the perception and processing of light and help to better identify problematic light levels. Eventually, dealing with ecological light pollution would ideally involve cooperation with physical scientists and engineers to help improve the equipment that can help to avoid ecological light pollution at a critical point in time for ecosystems. Such technical developments are then expected to help control, limit, or even stop the negative impact of light pollution.

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Authors’ contributions Ahmed AA Hussein: Conceptualization, funding acquisition, writing—original draft, writing—review and editing
Erik Bloem: Writing—original draft
István Fodor: Conceptualization, writing—original draft
El-Sayed Baz: Conceptualization, visualization, writing—original draft
Menerva M. Tadros: Conceptualization, writing—review and editing
Maha FM. Soliman: Conceptualization, supervision, visualization, writing—review and editing
Nahla S El-Shenawy: Conceptualization, supervision, visualization, writing—review and editing
Joris M. Koene: Conceptualization, supervision, funding acquisition, visualization, writing—original draft, writing—review and editing

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References

Abt KF, Schultz G (1995) Auswirkungen der Lichtemissionen einer Gro’s gewächshausanalyse auf den nächtlichen Vogelzug. Corax 16:17–29
Albertano P, Barsanti L, Passarelli V, Guaitieri P (2000) A complex photoreceptive structure in the cyanobacterium Leptolyngbya sp. Micron 31:27–34
Arendt D (2008) The evolution of cell types in animals: emerging principles from molecular studies. Nat Rev Genet 9:868–882
Arendt D, Tessmar K, de Campos-Baptista M-IM, Dorresteijn A, Wittbrodt J (2002) Development of pigment-cup eyes in the polychaete Platynereis dumerilii and evolutionary conservation of larval eyes in Bilateria. Development 129:1143–1154
Arendt D, Tessmar-Raible K, Snyman H, Dorresteijn AW, Wittbrodt J (2004) Ciliary photoreceptors with a vertebrate-type opsin in an invertebrate brain. Science 306:869–871
Badman DG (1966) Effects of light on the orientation of the snail Physa pomillia in a weak magnetic field. Psychol Forsch 29:360–371
Bailey SE (1981) Circannual and circadian rhythms in the snail Helix aspersa Müller and the photoperiodic control of annual activity and reproduction. Journal of Comparative Physiology A: Neuroethology, Sensory, Neural, and Behavioral Physiology 142:89–94
Baker J (1990) Toad aggregations under street lamps. British Herpetological Society Bulletin 31:26–27
Barré, K., Spoelstra, K., Bas, Y., Challeat, S., Kiri Ing, R., Azam, C., Zissis, G., Lapostolle, D., Kerbiriou, C., Le Viol, I., 2020. Artificial light may change flight patterns of bats near bridges along urban waterways. Anim Conserv acv.12635. https://doi.org/10.1111/acv.12635
Baz E-S, Wei H, Grosshans J, Stengl M (2013) Calcium responses of circadian pacemaker neurons of the cockroach Rhyparobia maderae to acetylcholine and histamine. J Comp Physiol A 199:365–374
Beckmann H, Hering L, Henze MJ, Kelber A, Stevenson PA, Mayer G (2015) Spectral sensitivity in Onychophora (velvet worms) revealed by electroretinograms, phototactic behaviour and opsin gene expression. J Exp Biol 218:915–922
Bedrosian TA, Fonken LK, Walton JC, Haim A, Nelson RJ (2011a) Dim light at night provokes depression-like behaviors and reduces CA1 dendritic spine density in female hamsters. Psychoneuroendocrinology 36:1062–1069
Bedrosian TA, Fonken LK, Walton JC, Nelson RJ (2011b) Chronic exposure to dim light at night suppresses immune responses in Siberian hamsters. Biol Lett 7:468–471
Benbellil-Tafoughalt, S., Sahoune, M., de Vauvleur, A., Moali, A., 2009. Effects of temperature and photoperiod on growth and reproduction of the land snail Helix aperta Born (Gastropoda, Pulmonata)
Berger D, Gotthard K (2008) Time stress, predation risk and diurnal–nocturnal foraging trade-offs in larval prey. Behav Ecol Sociobiol 62:1655–1663
Bobkova MV (1998) Structural and functional organization of the peripheral part of the visual system of the common pond snail Lymnaea stagnalis. J Evol Biochem Physiol 34:531–546
Bolhken S, Joose J (1981) The effect of photoperiod on female reproductive activity and growth of the freshwater pulmonate snail Lymnaea stagnalis kept under laboratory breeding conditions. International Journal of Invertebrate Reproduction 4:213–222. https://doi.org/10.1080/01651269.1981.10553430
Boldogh S, Dobrosi D, Samu P (2007) The effects of the illumination of buildings on house-dwelling bats and its conservation consequences. Acta Chiropterologica 9:527–534
Bouchet P, Rocroi J-P, Frýda J, Hausdorf B, Ponder W, Valdés Á, Warén A (2005). Classification and nomenclature of gastropod families
Bradshaw WE, Holzapfel CM (2010) Light, time, and the physiology of biotic response to rapid climate change in animals. Annu Rev Physiol 72:147–166
Brainard GC, Richardson BA, Petterborg LJ, Reiter RJ (1982) The effect of different light intensities on pineal melatonin content. Brain Res 233:75–81
Britt AB (1996) DNA damage and repair in plants. Annu Rev Plant Biol 47:75–100
Brown AM, Brown HM (1973) Light response of a giant Aplysia neuron. The Journal of general physiology 62:239–254
Bruce-White, C., Shardlow, M., 2011. A review of the impact of artificial night sky brightness. Mon Not R Astron Soc 328:689–707
Bruce-White, C., Shardlow, M., 2011. A review of the impact of artificial night sky brightness. Mon Not R Astron Soc 328:689–707
Carrascal LM, Santos T, Telleria JL (2012) Does day length affect winter bird distribution? Testing the role of an elusive variable. PLoS One 7:e32733
Chase RB (2002) Behavior & its neural control in gastropod molluscs. Oxford University Press on Demand
Cinzano P, Falchi F, Elvidge CD (2001) The first world atlas of the night sky. Oxford University Press
Cos S, Mediavilla D, Martínez-Campa C, González A, Alonso-González C, Sánchez-Barceló EJ (2006) Exposure to light-at-night increases the growth of DMBA-induced mammary adenocarcinomas in rats. Carcinogenesis 27:235–240
Cronin TW, Johnsen S, Marshall NJ, Warrant EJ (2014). Visual ecology. Princeton University Press
Dauchy RT, Sauer LA, Blask DE, Vaughan GM (1997) Light contamination during the dark phase in “photorhythmically controlled” animal rooms: effect on tumor growth and metabolism in rats. Comparative Medicine 47:511–518
Davies TW, Duffy JP, Bennie J, Gaston KJ (2014) The nature, extent, and ecological implications of marine light pollution. Front Ecol Environ 12:347–355
De Molenaar JG, Jonkers DA, Henkens R (1997) Wegeverlichting en natuur. I. Een literatuurstudie naar de werking en effecten van licht en verlichting op de natuur
De Molenaar JG, Jonkers DA, Sanders ME (2000) Road illumination and nature; III local influence of road lights on a black-tailed godwit (Limosa l. limosa) population
Del Pilar Gomez M, Angueyra JM, Nasi E (2009) Light-transduction in algal palatability and herbivore feeding behavior and performance. Oecologia 180:453–462
Doring CC, Kumar S, Tumu SC, Kourtessis I, Hausen H (2020) The visual pigment xenopsin is widespread in protostome eyes and impacts the view on eye evolution. eLife 9:e55193
Duarte C, López J, Benítez S, Manriquez PH, Navarro JM, Bonta CC, Torres R, Quijón P (2016) Ocean acidification induces changes in algal palatability and herbivore feeding behavior and performance. Oecologia 180:453–462
Eisenbeis G, Hassel F (2000) Zur Anziehung nachtaktiver Insekten durch Strassenlaternen. Natur und Landschaft 75:145–156
Evans Ogden LJ (1996) Collision course: the hazards of lighted structures and windows to migrating birds Fatal Light Awareness Program (FLAP):3
Evans JA, Elliott JA, Gorman MR (2007) Circadian effects of light no brighter than moonlight. J Biol Rhythm 22:356–367
Fain GL, Hardie R, Laughlin SB (2010) Phototransduction and the evolution of photoreceptors. Curr Biol 20:R114–R124
Falkenberg JC, Clarke JA (1998) Microhabitat use of deer mice: effects of interspecific interaction risks. J Mammal 79:558–565
Fanini L, Hughes LE, Springthorpe R, Tosetto L, Lowry JK (2016) Surface activity patterns of macrofauna on pocket, tidal beaches: insights into the role of wrack and artificial lighting. Reg Stud Mar Sci 7:63–71
Fodor I, Hussein AA, Benjamin PR, Koene JM, Pirger Z (2020) The unlimited potential of the great pond snail, Lymnaea stagnalis. eLife 9:e56962. https://doi.org/10.7554/eLife.56962
Foster RG, Rogenneben T (2008) Human responses to the geophysical daily, annual and lunar cycles. Curr Biol 18:R784–R794
Frank KD (1988) Impact of outdoor lighting on moths: an assessment. Journal of the Lepidopterists’ Society (USA)
Frank KD, Rich C, Longcore T (2006) Consequences of artificial night lighting
Gaston KJ, Bennie J (2014) Demographic effects of artificial nighttime lighting on animal populations. Environ Rev 22:323–330
Gaston KJ, Bennie J, Davies TW, Hopkins J (2013) The ecological impacts of nighttime light pollution: a mechanistic appraisal. Biol Rev 88:912–927
Gaynor KM, Hojnowski CE, Carter NH, Brashears JS (2018) The influence of human disturbance on wildlife nocturnality. Science 360:1232–1235. https://doi.org/10.1126/science.aar7121
Gehring WJ (2014) The evolution of vision. Wiley Interdiscip Rev Dev Biol 3:1–40. https://doi.org/10.1002/wdev.96
Gehring W, Seimiya M (2010) Eye evolution and the origin of Darwin’s eye prototype. Italian Journal of Zoology 77:124–136. https://doi.org/10.1080/11250001003795350
Gerrish GA, Morin JG, Rivers TJ, Patrawala Z (2009) Darkness as an ecological resource: the role of light in partitioning the nocturnal niche. Oecologia 160:525–536
Giacconi C (2006) Efecto de la contaminación lumínica sobre la abundancia, riqueza y comportamiento de la macrofauna de playas arenosas de la IV Región. Centro de Egresados de Ingeniería en Recursos Naturales Renovables, Memorias y Publicaciones, Universidad de Chile. Resumen
Gornot A (1990) Photoperiod and temperature interaction in the determination of reproduction of the edible snail, Helix pomatia. J Reprod Fertil 90:581–585
González SA, Yáñez-Navea K, Muñoz M (2014) Effect of coastal urbanization on sandy beach coleoptera Phaleria maculata (Kulzer, 1959) in northern Chile. Mar Pollut Bull 83:265–274
Gorman MR, Kendall M, Elliott JA (2005) Scopitic illumination enhances entrainment of circadian rhythms to lengths light: dark cycles. J Biol Rhythm 20:38–48
Gotow T, Nishi T (2009) A new photosensory function for simple photoreceptors, the intrinsically photosensitive neurons of the sea slug Onchidium. Front Cell Neurosci 3:18
Ramirez MD, Pairett AN, Pankey MS, Serb JM, Speiser DI, Swafford AJ, Rand AS, Bridarolli ME, Dries L, Ryan MJ (1997) Light levels influence circadian mechanisms in the context of integrative photoreceptor cell biology. Vis Neurosci 28:265–279

Ramirez MD, Pairatt AN, Pankey MS, Serb JM, Speiser DI, Swafford AJ, Oakley TH (2016) The last common ancestor of most bilaterian animals possessed at least nine opsins. Genome biology and evolution 8:3640–3652

Rand AS, Bridarolli ME, Dries L, Ryan MJ (1997) Light levels influence female choice in túngara frogs: predation risk assessment? Copeia 1997:447–450

Rawlinson KA, Lapraz F, Ballister ER, Terasaki M, Rodgers J, McDowell RJ, Girstmair J, Criswell KE, Boldogkoi M, Simpson F (2019) Extraocular, rod-like photoreceptors in a flatworm express xenopsin photopigment. ELife 8:e45465

Reiter RJ, Tan DX, Sanchez-Barcelo E, Mediavilla MD, Gitto E, Korkmaz A (2011) Circadian mechanisms in the regulation of melatonin synthesis: disruption with light at night and the pathophysiological consequences. J Exp Integr Med 1:13–22

Ricciardi A, Rasmussen JB (1998) Predicting the identity and impact of future biological invaders: a priority for aquatic resource management. Can J Fish Aquat Sci 55:1759–1765

Rossetti Y, Cabanac M (2006) Light versus temperature: an intersensitivity conflict in a gastropod (Lymnaea auricularia). J Therm Biol 31:514–520

Rumpho ME, Worful JM, Lee J, Kannan K, Tyler MS, Bhattacharya D, Moustafa A, Manhart JR (2008) Horizontal gene transfer of the algal nuclear gene psbO to the photosynthetic sea slug Elysia chlorotica. Proc Natl Acad Sci 105:17867–17871. https://doi.org/10.1073/pnas.0804968105

Rydell J (1992) Exploitation of insects around streetlamps by bats in Sweden. Funct Ecol 6:744–750

Sakikabara M, Aritaka T, Iizuka A, Suzuki H, Horikoshi T, Lukowiak K (2005) Electrophysiological responses to light of neurons in the eye and statocyst of Lymnaea stagnalis. J Neurophysiol 93:493–507. https://doi.org/10.1152/jn.00692.2004

Salmon M, Tolbert MG, Painter DP, Goff M, Reiners R (1995) Behavior of loggerhead sea turtles on an urban beach. II Hatching orientation. Journal of Herpetology 58:576–578

Sarfati R, Hayes J, Sarfati É, Peleg O (2020) Spatiotemporal reconstruction of emergent flash synchronization in firefly swarms via stereoscopic 360-degree cameras. bioRxiv

Schwartz A, Henderson RW (1991). Amphibians and reptiles of the West Indies: descriptions, distributions, and natural history. University Press of Florida

Shubin N, Tabin C, Carroll S (2009) Deep homology and the origins of evolutionary novelty. Nature 457:818–823

Sinha RP, Håder D-P (2002) UV-Induced DNA damage and repair: a review. Photochemical & Photobiological Sciences 1:225–236

Sokolove PG, McCrone EJ (1978) Reproductive maturation in the slug, Limax maximus, and the effects of artificial photoperiod. J Comp Physiol 125:317–325

Stephens CJ, Stephens GC (1966) Photoperiodic stimulation of egg laying in the land snail Helix aspersa. Nature 212:1582

Stevens RG (2009) Light-at-night, circadian disruption and breast cancer: assessment of existing evidence. Int J Epidemiol 38:963–970

Stoll CJ, Bijlsma A (1973) Optic-nerve responses in Lymnaea-Stagnalis (l(Pulmonata, Basommatophora)) to photic stimulation of eye. Proceedings of The Koninklijke Nederlandse Akademie van Wetenschappen Series C-Biological and Medical Sciences 76:406

Summers CG (1997) Phototactic behavior of Bemisia argentifolii (Homoptera: Aleyrodidae) crawlers. Ann Entomol Soc Am 90:372–379

Takigami S, Sunada H, Horikoshi T, Sakakibara M (2014) Morphological and physiological characteristics of dorsal photoreceptors in Lymnaea stagnalis. Biophysics 10:77–88. https://doi.org/10.2142/biophysics.10.77

Ter Maat A, Zonneveld C, de Visser JAG, Jansen RF, Montagne-Wajer K, Koene JM (2007) Food intake, growth, and reproduction as affected by day length and food availability in the pond snail Lymnaea stagnalis. Am Malacol Bull 23:113–120

Ter Maat A, Pienneman AW, Koene JM (2012) The effect of light on induced egg laying in the simultaneous hermaphrodite Lymnaea stagnalis. J Molluscan Stud 78:262–267

Udaka H, Goto SG, Numata H (2008) Effects of photoperiod and acclimation temperature on heat and cold tolerance in the terrestrial slug, Lehmannia valeta (Pulmonata: Limacidae). Appl Entomol Zool 43:547–551

Underwood CN, Davies TW, Queirós AM (2017) Artificial light at night alters trophic interactions of intertidal invertebrates. J Anim Ecol 86:781–789. https://doi.org/10.1111/1365-2656.12670

van Duivenboden YA (1982) Non-ocular photoreceptors and phototaxis in the pond snail Lymnaea stagnalis (L.). J Comp Physiol 149:363–368

Vasquez RA (1994) Assessment of predation risk via illumination level: facultative central place foraging in the cricetid rodent Phyllotis darwini. Behav Ecol Sociobiol 34:375–381

Velaere RD, Sauer CD, Walden KK, Fahrbach SE, Robertson HM (2005) Pteropsin: a vertebrate-like non-visual opsin expressed in the honey bee brain. Insect Biochem Mol Biol 35:1367–1377

Verheijen FJ (1960) The mechanisms of the trapping effect of artificial light sources upon animals. Arch Néerl Zool 13:1

Verheijen FJ (1985) Photopollution: artificial light optic spatial control of emergent flash synchronization in firefly swarms via stereoscopic 360-degree cameras. Elife 6:e23435

Volkensky P, Pergner J, Liegertova M, Benito-Gutierrez E, Arendt D, Nurius P, Matic K, Koene JM (2007) Food intake, growth, and reproduction as affected by day length and food availability in the pond snail Lymnaea stagnalis. Am Malacol Bull 23:113–120

Wayne NL, Block GD (1992) Effects of photoperiod and temperature on egg-laying behavior in a marine mollusc, Aplysia californica. Biol Bull 182:8–14

Wensel TG (2008) Signal transducing membrane complexes of photoreceptor outer segments. Vis Res 48:2052–2061. https://doi.org/10.1016/j.visres.2008.03.010

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