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GCB REVIEW



Impacts of artificial light at night in marine ecosystems—A review

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Abstract

The globally widespread adoption of Artificial Light at Night (ALAN) began in the mid-20th century. Yet, it is only in the last decade that a renewed research focus has emerged into its impacts on ecological and biological processes in the marine environment that are guided by natural intensities, moon phase, natural light and dark cycles and daily light spectra alterations. The field has diversified rapidly from one restricted to impacts on a handful of vertebrates, to one in which impacts have been quantified across a broad array of marine and coastal habitats and species. Here, we review the current understanding of ALAN impacts in diverse marine ecosystems. The review presents the current state of knowledge across key marine and coastal ecosystems (sandy and rocky shores, coral reefs and pelagic) and taxa (birds and sea turtles), introducing how ALAN can mask seabird and sea turtle navigation, cause changes in animals predation patterns and failure of coral spawning synchronization, as well as inhibition of zooplankton Diel Vertical Migration. Mitigation measures are recommended, however, while strategies for mitigation were easily identified, barriers

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to implementation are poorly understood. Finally, we point out knowledge gaps that if addressed would aid in the prediction and mitigation of ALAN impacts in the marine realm.

KEYWORDS

artificial light at night (ALAN), conservation guidelines, coral reefs, marine ecosystem, pelagic organisms, rocky intertidal shores, sandy beach, seabirds, sea-turtles

1 | INTRODUCTION

Artificial Light at Night (ALAN) is a widespread, pervasive, and expanding form of pollution (Gaston et al., 2021) that has come to be recognized as a major 21st century global change issue (Davies & Smyth, 2018). Its impacts span the biological hierarchy ranging from those on organism physiology through to changes in the composition of ecological communities (Sanders et al., 2021). It is now broadly accepted that ALAN has been reshaping nature for more than a century.

Although research into the prevalence and impacts of ALAN in marine ecosystems has somewhat lagged behind terrestrial, the last 5 years have seen a dramatic advance in our understanding. We now know that at least 22% of coastal regions are exposed to ALAN (Davies et al., 2014), and the light from cities is sufficient to elicit biological responses in animals on the seafloor in adjacent habitats (Ayalon, Rosenberg, et al., 2021; Davies et al., 2020). 1.9 million km² of the world's coastal seas are exposed to ALAN at 1 m depth, 1.6 million km² at 10 m depth, and 840,000 km² at 20 m depth (Figure 1a) (Smyth et al., 2021). The most exposed regions include the Mediterranean (Figure 1b), the Red Sea and Persian Gulf (Figure 1c), and the seas of South-East Asia (Figure 1d). ALAN is even prevalent across those areas of the ocean most valued by humanity, with 20% of the world's contiguous Marine Protected Areas exposed across 100% of their range (Davies et al., 2016).

The potential for ALAN to impact the wide array of organisms, processes, and habitats in the sea for which light cycles are critical had remained largely unexplored until recently (Davies et al., 2014; Longcore & Rich, 2004). These include: diel vertical migrations (Berge et al., 2020)—the largest migration of biomass on the planet (Hayes, 2003); coral spawning (Ayalon, Rosenberg, et al., 2021), and symbiosis (Ayalon, Benichou, et al., 2021)-which are key for the maintenance of coral reefs; consumer-resource interactions (Bolton et al., 2017; Maggi, Bongiorni, et al., 2020; Underwood et al., 2017) that are known to drive top down structuring of marine ecosystems (Paine, 1966); migrations and orientation of marine organismscritical for their survival (Navarro-Barranco & Hughes, 2015; Torres et al., 2020); and the recruitment of sessile invertebrate larvae into marine habitats (Davies et al., 2015; Lynn, Quintanilla-Ahumada, et al., 2021), (Figure 2). All these processes are fundamental to the health of marine ecosystems, and all are known to depend on the cycles, spectra or intensity of sun or moonlight.

Here, we bring together recognized experts in marine ALAN across habitats and taxonomic groups to review these recent

advances, with the aim of providing a gateway to research in the field. First, we review progress in sandy beaches, rocky intertidal shores, shallow water coral reefs, and pelagic environments (Figure 2). Second, we evaluate the current state of litigation and management options available to conservation practitioners. Finally, we identify key knowledge gaps and highlight key questions for future research.

2 | BIOLOGICAL EFFECTS OF ALAN IN THE MARINE BIOTA

In this section we have compiled the most relevant information obtained on key species belonging to different marine habitats across sandy beaches (including sea turtles), rocky intertidal shores, shallow water coral reefs, and pelagic environments. A special section was created for seabirds due to their high mobility and, therefore, presence in different marine realms. Despite the research gaps and limitations-to name a few, for example, lack of multistressor experiments including ALAN as a factor, and difficulties to acquire data to define more precisely ecological relevant light intensities to be tested (see Box 1) (Aulsebrook et al., 2022)-ALAN can be recognized as major sensory pollutant of concern due it is obvious and widespread effects on pathways associated with natural circadian regulations in the marine biota. It is also important to note that experimental approaches using high ALAN levels (not considered environmental realistic) in many of the short-term experiments presented here consist in an important step to understanding the mechanisms and long-term effects of the chronic disturbance caused by ALAN.

2.1 | Sandy beach ecosystems

Many sandy beach species are known for their day/ night activity rhythms controlled by natural light cycles. Consequently, the dramatic ongoing expansion of ALAN in these ecosystems is expected to exert a significant effect on sandy beach biodiversity (Figure 3).

Exposed sandy beach ecosystems represent over 80% of the ice-free coastline (Bascom, 1980). These ecosystems are considered highly valuable from an ecologic, economic and cultural points of view (King & Symes, 2004; McLachlan et al., 2013; Pendleton et al., 2006). However, the growing urbanization and the increase of

FIGURE 1 The depth of biologically important artificial light at night (ALAN) (*Zc*): around the world's coastlines (a); in the Mediterranean and Northeast Atlantic (b); in the Black Sea, the Red Sea and the Persian Gulf (c); and in the Gulf of Thailand, Andaman Sea, South China Sea and the Java Sea (d). The legend inset details the depths (m) to which biologically important ALAN penetrates the sea. The data is derived from the relationship between The New World Atlas of Artificial Night Sky Brightness (Falchi et al., 2016) and sea surface irradiances (Davies et al., 2020), coupled with the monthly climatologies of globally inherent optical water properties and validated against in situ data collected from the Persian Gulf (Tamir et al., 2017). "Biologically important ALAN" is defined as the minimum irradiances of white light that elicit diel vertical migration in female *Calanus* copepods (Båtnes et al., 2013) [see Smyth et al., 2021 for further details]. Maps are representative of average ALAN penetration over a typical year. Full dataset is available to download from https://doi.pangaea.de/10.1594/PANGA EA.929749.

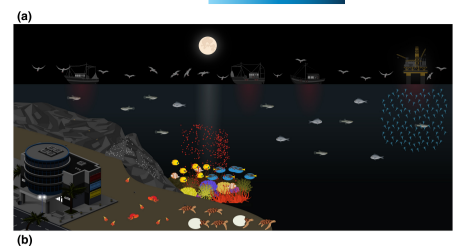
human population near sandy shores has become a threat to these ecosystems (Jaramillo et al., 2021). One of the most important environmental stressors associated with the urbanization of sandy beaches is ALAN (Gonzalez et al., 2014; Schlacher et al., 2016).

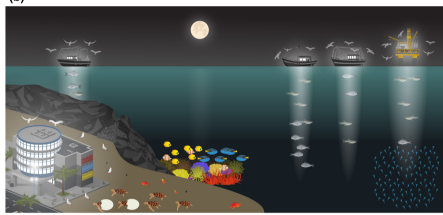
Sandy beaches support an abundant and diverse fauna, being crustaceans (mainly talitrid amphipods, cirolanid and oniscoid isopods and hippid crabs), polychaetes and bivalves the main taxonomic groups (Brown & McLachlan, 1990; Jaramillo et al., 2003). Several sandy beach species such as amphipods, isopods, and insects are known for their dynamic day/night activity rhythms (Fallaci et al., 1999, 2002; Jaramillo et al., 2003; Kennedy et al., 2000), which are controlled by natural geophysical cycles of day and night, and moonlight (Jaramillo et al., 2003; Meschini et al., 2008; Ugolini et al., 2003). For example, these invertebrates rely on darkness to use natural light signals and return to their burrows after down shore migrations (Fallaci et al., 2002). Consequently, the ongoing expansion of ALAN in these ecosystems is expected to exert both

a significant and negative effect on sandy beach biodiversity (e.g., Duarte et al., 2019; Luarte et al., 2016). However, despite growing concerns on the impacts of ALAN, studies evaluating its effects on sandy beach organisms, other than sea turtles (e.g., Rivas et al., 2015; Dimitriadis et al., 2018, see specific section 2.5. for this group), remain very scarce (e.g., Lynn, Flynn, et al., 2021; Quintanilla-Ahumada et al., 2021). The studies carried out so far have evaluated the effects of ALAN on locomotor activity, feeding rates, absorption efficiency and growth rate, physiology, distribution, and abundance of different species of invertebrates (see below).

One of the first studies evaluating the ALAN effects on locomotor activity of sandy beach animals was conducted by Bregazzi and Naylor (1972). Those authors showed that in laboratory conditions ALAN (~200 lux) almost entirely inhibited the locomotor activity of the talitrid amphipod *Talitrus saltator*. Interestingly, these individuals recovered their locomotor activity pattern immediately after the light was removed. Similar results were registered by Luarte

FIGURE 2 (a) Different marine environments not affected by Artificial Light Pollution at Night (ALAN), and (b) marine environments under the potential impacts of ALAN: (i) Sandy beaches effects on invertebrate species day-night activity rhythms and biodiversity, effects in the on-beach orientation of adults and hatchling turtles, and seabirds fledgling grounded by ALAN; (ii) Rocky intertidal shores - influence in metabolic activity/ behavior of primary producers, sessile and mobile animals; (iii) Shallow water coral reefs - effects on gametogenesis and the synchronization of gamete release in prominent coral species, and negative impacts over fish reproduction, (iv) Pelagic environment - inhibition of vertically migrating zooplankton, and disorientation and mortality of seabirds.





et al. (2016), who using laboratory experiments showed that the locomotor activity of the talitrid amphipod Orchestoidea tuberculata was significantly reduced in the presence of ALAN (60 lux). Similarly. Duarte et al. (2019) found that in laboratory conditions (120 lux), ALAN reduced the locomotor activity of the oniscoid isopod Tylus spinulosus. More recently, Lynn et al. (2021,b) also found that ALAN (80 lux) disrupted the locomotor activity of the talitrid amphipod Americorchestia longicornis. Consistent with the results reported by Bregazzi and Naylor (1972), A. longicornis was found to recover its natural rhythm of activity shortly after ALAN was removed from the system (Lynn, Flynn, et al., 2021). By contrast, Fanini et al. (2016) found that the locomotor activity of the amphipod Platorchestia smithi was similar in a beach exposed to ALAN with respect to another that was not exposed. However, this last study was correlational and should be considered with caution. More recently 0.2 lux white lighting lower in brightness than a full moon (equivalent to artificial sky glow) has been demonstrated to reduce locomotor activity and disorientate the migration behavior of Talitrus saltator (Torres et al., 2020).

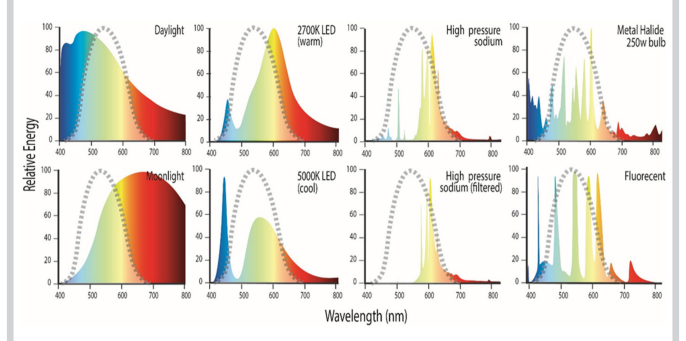
Only a scarce number of studies have evaluated ALAN effects on aspects such as feeding rates, absorption efficiency and growth rates (Luarte et al., 2016; Lynn, Flynn, et al., 2021; Quintanilla-Ahumada et al., 2022). The presence of ALAN reduced the consumption rate, absorption efficiency and growth rate in the amphipods O. tuberculata (Luarte et al., 2016) and A. longicornis (Lynn, Flynn, et al., 2021), meanwhile ALAN (from 0 to

100 lx) did not affect T. spinulosus's growth rate but increased its absorption efficiency (Quintanilla-Ahumada et al., 2022). The absorption efficiency results should be considered with precaution. because of methodological restrictions, as the animals had to be maintained without sand, which, eventually, could modify the ALAN effects on this biological trait). The RNA: DNA ratio is a relatively new indicator of the physiological or nutritional condition of organisms (Buckley et al., 1999; Chícharo & Chícharo, 2008). Recent studies by Quintanilla-Ahumada et al. (2021, 2022), used this molecular tool to evaluate ALAN effects in sandy beach ecosystems, found that RNA:DNA ratio in the insect Phalerisida maculata and in the isopod T. spinulosus, declined in the presence of ALAN, indicating detrimental physiological effects. ALAN also shows important effects on the abundance and distribution of sandy beach organisms. Gonzalez et al. (2014) applied correlative analyses and found that the abundance of the insect P. maculata was negatively related with the night sky quality (an indirect indicator of ALAN). Duarte et al. (2019) registered that the distribution of T. spinulosus was modified by ALAN (120 lux and less), with individuals avoiding the lit areas and therefore restricting their habitat availability. Importantly, ALAN effects decreased with increasing distance from the light sources (Duarte et al., 2019). Despite the work conducted this far, direct evidence of ALAN effects on sandy beach organisms remains limited only to studies focusing on this stressor upon single species (e.g., Duarte et al., 2019; Luarte et al., 2016).

BOX 1 Challenges in measuring artificial light in biological studies

Measuring light for ecological studies is still undefined and is poorly understood by biologists who have used a wide range of techniques and instruments to measure light at scales ranging from a few centimeters to tens of kilometers. The wide range of instruments employed typically use different measurement systems and units, which means study results cannot be confidently compared. There is currently no globally recognized standard method, or agreed unit of measurement, for monitoring biologically active light (Barentine, 2019).

The figure below demonstrates this. Commercial lux meters only quantify light that is within the CIE curve (area under grey dashed line) and exclude light wavelengths that fall outside of the CIE curve.



Credit: Kellie Pendoley.

Visible light can be described by two physical parameters: wavelength and intensity. The relative distribution and weighting of different wavelengths in emitted light determines its color; however, how this light is perceived by the observer is also a function of the physiology of the receiving detector (e.g., eye). The intensity of light reaching a detector is a function of distance from the light source as light waves spread out from the light source the number of waves falling on a defined area decreases proportional to the distance travelled. Light is measured either radiometrically or photometrically. Radiometry is the measurement of wavelengths across the entire electromagnetic spectrum. In biological applications this is typically restricted to the ultraviolet, visible, and infrared region of the spectrum between 350 and 800 nm and is measured in watts per meter squared. Most commercial light measurement equipment records photometric light which is visible light wavelengths weighted specifically to the sensitivity of the human eye (CIE, 2005) and is reported in units of lumens per meter squared or Lux. Photometric detectors have reduced sensitivity to wavelengths below 450 nm (blue light) or above 650 nm (red light). Consequently, photometric instruments commonly used to quantify light in biological studies, such as lux meters, do not account for the blue light that is most visible to biological receptors.

Lux meters are further limited in that they were designed for use in measuring light in buildings, have poor sensitivity to low light levels and cannot detect sky glow or dim light in a field setting. Finally, they do not provide any spectral information, i.e., the relative distribution of light wavelengths, in the light source.

The limitations and challenges around measuring ecological light on scales ranging from landscape to small scale is discussed in more detail in Commonwealth (2020); Longcore et al. (2020), Hänel et al. (2018), Jechow and Hölker (2019) and Jechow et al. (2019).

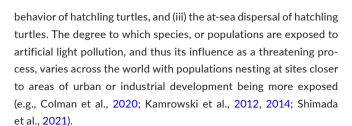
It is important to note that the light intensities used in some of these studies were those recorded under the light source (mainly in Chilean sandy beaches) in the promenade area, located very close to the beach. Such light intensities could be higher than those occurring in the intertidal area, mainly in the middle and lower intertidal zone. Therefore, future studies should consider using the light intensity

directly recorded at the intertidal zone. However, at least for the Chilean coast, in extreme heavy polluted beaches, light intensity values in the intertidal zone can be as high or higher than those used in these studies, mainly in the upper intertidal zone (Duarte unpublished data). Another important consideration should be to expand ALAN studies to beaches located in different geographic areas, for example tropical ones, which with the exception of sea turtle species, have not received special attention regarding to ALAN.

2.1.1 Sea-Turtles

It is well established that ALAN, even at low levels, is a threat influencing all seven species of marine turtle, primarily as hatchlings or nesting adult females (Salmon, 2003; Witherington & Bjorndal, 1991a, 1991b) (Figure 4). Artificial light pollution is known to influence (i) the on-beach orientation and nest site selection of adult female turtles, (ii) the on-beach orientation and sea finding

FIGURE 3 Los Choros Sandy beach, coast of Coquimbo, North of Chile (Credit: Josué Navarrete).



Artificial light primarily impacts marine turtles during nesting or hatchling life stages, and consequently it is predominantly linked to early life stage mortality. Thus, it could, along with other threats, cause gradual decline in the reproductive output of a nesting area. However, while there are several studies examining beach specific exposure, or short-term (season) impacts, quantifying the long-term impacts of artificial light pollution-caused mortality are difficult because marine turtles can take decades to reach maturity, and impacts of light pollution contribute to other pressures over turtles' lifetimes to influence population viability. Hence, understanding the degree to which nesting sites for each population of turtles are





FIGURE 4 Sea turtles hatching in Heron Island, Australia (Credit; Levy, O).

exposed to artificial light pollution is important, so that site-specific interventions can be implemented.

Typically, female turtles lay their clutches above the highwater mark, and often in the primary dune systems or high up on the beach. Hatchlings generally emerge from nests at night and orient themselves from the nest site to the ocean, ideally as fast as possible (Wyneken & Salmon, 1992). After emerging they generally show a preference of moving towards horizons which are low and bright, and moving away from horizons, which are dark and elevated (e.g., Limpus & Kamrowski, 2013; Lucas et al., 1992; Pendoley & Kamrowski, 2015; Salmon et al., 1995) and using these cues they can navigate across the beach to the water. Exposure to coastal light pollution disrupts the natural orientation cues and leads to the disorientation and misorientation of hatchlings because lights obscure the natural horizons (Witherington & Bjorndal, 1991a, 1991b).

Hatchling sea finding ability is influenced by both the wavelength and intensity of artificial light (Cruz et al., 2018; Witherington & Bjorndal, 1991b). Such ability is significantly compromised by exposure to shorter wavelength lights, even at lower intensity (Celano et al., 2018; Salmon, 2003). Importantly, it is becoming clear that the thresholds of concern for both wavelength and intensity of artificial light are likely to vary within and among species (Fritsches, 2012). The impacts of exposure can be influenced by the presence or absence of other natural (dune height and structure, vegetation) or unnatural cues (presence of buildings or artificial structures) (Kamrowski et al., 2015; Salmon, 2003), highlighting the need for site-specific research on orientation thresholds and light-reduction interventions.

Once at sea, hatchling turtles will swim actively for around 24 to 48 h (Wyneken & Salmon, 1992). This period, known as the swim frenzy, enables hatchlings to move quickly from nearshore to offshore waters (Wyneken & Salmon, 1992). During the swim frenzy the hatchlings are using multiple cues to enable their directional swimming - these include swimming towards the low, light, horizon, and swimming perpendicular to wave fronts (Lohmann et al., 2017; Salmon & Wyneken, 1987; Wilson et al., 2021). According to Salmon and Wyneken (1987) light cues are important for at sea dispersal, however, there is likely to be a distance offshore where the cue is either not available or not used. Although this distance is currently unknown, there is a growing empirical basis demonstrating that offshore dispersal for marine turtle hatchlings is compromised by light pollution originating from landbased or marine structures, such as infrastructure like jetties (Cruz et al., 2018; Truscott et al., 2017; Wilson et al., 2018). Continuing to advance knowledge on how the at sea dispersal is influenced by artificial lights from shore, or offshore infrastructure is a key avenue for further research as human developments expand along

For nesting turtles there has been substantial research on factors that influence nest site selection (i.e., the placement of clutches on a beach). Factors including distance from vegetation (most nests being laid closer to the vegetation line—Hays et al., 1995;

Kelly et al., 2017), elevation and beach slope (Patrício et al., 2018; Wood & Bjorndal, 2000) and exposure to artificial light pollution (Salmon, 2003; Windle et al., 2018) have all been associated with nest site selection. There is also variation among species and region as to the relative importance of each. However, less research has been conducted to examine the influence of light pollution on nest site selection by nesting turtles. Among the studies, Salmon (2003) used data from a main nesting area in Florida to test nest site selection in relation to the degree females were exposed to artificial light spill onto beaches; and Windle et al. (2018) used a combination of light pollution data and turtle density data to examine the influence of artificial light pollution on nest site selection. Both studies concluded that turtles use darker beaches and select darker sections of beaches.

2.2 | Rocky intertidal shores

On a global scale, rocky intertidal shores are inhabited by diverse assemblages largely influenced by light for metabolic activities and behaviors at different life stages. It is reasonable to expect that the presence of ALAN may influence microphytobenthic and macroalgal primary producers, as well as sessile and mobile animals (Figure 5).

Intertidal habitats represent a thin line demarcating the boundary between land and marine masses. Among intertidal environments, rocky shores are characterized by large variations in abiotic conditions, including strong gradients in wave exposure, temperature, and desiccation. Despite the importance of such extreme factors, biotic interactions have shown strong determinants of distribution patterns and abundance of organisms. These features have historically made rocky shore intertidal habitats a natural laboratory to explore physiological and ecological processes and mechanisms, which have been responsible for key conceptual advancements on ecosystem functioning (Menge & Branch, 2001). ALAN, however, represents a guite novel and peculiar source of disturbance, whose effects are hardly predictable from knowledge on other stressors. In fact, it is expected to act on pathways associated with natural circadian regulations, which are related to different light optima and sensitivities among species and life stages (Davies & Smyth, 2018). Although to date the literature on effects of ALAN on intertidal rocky shores is not particularly numerous, effects at the scale of both the individual and the ecosystem have already been highlighted.

Intertidal rocky shores host a diverse array of organisms, which can be primarily categorized into sessile and vagile. Sessile species are those most diverse in terms of both size, taxonomy and trophic category. A large percentage of these species belongs to microbial biofilm. Despite low visibility, its autotrophic component microphytobenthos, MPB) represents one of the main groups of primary producers in intertidal habitats and, in association with heterotrophic microorganisms, a source of food for a plethora of grazers species (Jenkins & Hartnoll, 2001; Nagarkar et al., 2004). Seaweeds are the



macroscopic primary producers on intertidal rocky shores, sometimes present as dense macroalgal beds able to modify the abiotic conditions on the shore through their canopies. Sessile species also include a variety of animals, such as bivalves, barnacles, ascidians, bryozoans, hydrozoans, gastropods and polychaetes, among others. Mobile animals are abundant and diversified as well: these include either herbivores, carnivores and omnivores almost freely living on the different vertical portions of the intertidal habitats (e.g., limpets, gastropods, crabs, sea stars, sea urchins, small fish), or smaller species more strictly associated with larger ones, such amphipods, isopods and crustaceans.

In the microtidal Mediterranean system, where intertidal biofilm is dominated by bacteria, the presence of lit areas at night at intensities typically found along the coast (27 lux) was able to increase the biomass of MPB (here dominated by cyanobacteria) and its photosynthetic efficiency (Maggi & Benedetti-Cecchi, 2018). Subsequent studies indicated that ALAN effect was likely mediated by temporal changes in composition of mature assemblages (Maggi, Bertocci, & Benedetti-Cecchi, 2020; Maggi, Bongiorni, et al., 2020) and possibly related to different light optima among (groups of) species. Potential consequences of effects on autotrophic microorganisms are not limited to net oxygen emissions and indirect impacts on their consumers. In fact, the composition of microbial biofilm plays a key role in the settlement of larval stages of invertebrates and spores (Keough & Raimondi, 1995; Qian et al., 2007). Indeed, the first study investigating the role of ALAN (either 30 or 19 lux) for intertidal organisms focused on sessile larval stages of animals, revealing variable effects on settlement rates on PVC panels attached on wooden floating rafts in the UK (Davies et al., 2015). Results showed that 39% of analyzed taxa were influenced by ALAN, either positively or negatively; it was not surprising, given the importance of natural light as a cue for guiding larval recruitment and later survival (Mundy

& Babcock, 1998; Thorson, 1964). More recently, negative effects of ALAN on late settlers of barnacles have been documented both along natural shores in Chile (97-11 lux) (Notochthamalus scabrosus and Jehlius cirratus; Manríguez, Jara, et al., 2021; Manríguez, Quijon, et al., 2021) and in the North Atlantic on man-made structures (212-11 lux) (Semibalanus balanoides; Lynn, Flynn, et al., 2021); in these studies, lack of effects on early settlers suggested that, in presence of lit nights, metamorphosis was delayed or early mortality was increased in comparison to natural dark conditions.

As for sessile intertidal species, a big knowledge gap on effects on autotrophs still exists, with a complete lack of studies on macroalgal species. In this case, it is worth mentioning that ALAN effects could also influence non-trophic interactions mediated by algal canopies, such as facilitative effects exerted through the reduction of artificial light intensities for understory assemblages (Bruno & Bertness, 2001). In addition to sessile species or life stages, intertidal rocky reefs are inhabited by mobile individuals. Mobility is a great advantage in a habitat characterized by high abiotic variation. For example, many organisms have evolved predominantly nocturnal behaviors to avoid energy expenditure related to thermal stress or to reduce the risk of predation linked to visual stimuli (Manríquez et al., 2009; Wells, 1980). Different organs are involved in the perception of circadian changes in light intensity and of prey or predators, from relatively simple photoreceptors capable of forming sharp images in air in gastropods (Newell, 1965), to proper eyes in fish. Along the Chilean coasts, the abalone Concholepas concholepas, or "Loco," is an ecologically and economically key species. Like most mollusks, it uses both chemical and visual stimuli during sensory perception (Domenici et al., 2017; Manríquez et al., 2014). Chemoreception of odor cues through the osphradium is the main tool to monitor the presence of food items or predators; while information detected through its simple eyes modulates phototaxis

Like some invertebrate species, many fish species have evolved endogenous clock systems which regulate tidally and circadian organized behavioral rhythms (Helfman et al., 2009). Among them is the "Baunco", the rockfish Girella laevifrons, one of the most abundant fish in the littoral zone of Southeastern Pacific. A recent study showed an increase in oxygen consumption and activity of this fish under ALAN conditions (70 lux). Importantly, loss of a dark night period was able to modify or even stop the daily peak of activity of the Baunco, posing serious questions about the sustainability of intertidal fish populations in urban areas (Pulgar et al., 2019). Results are in accordance with those obtained on fish by manipulating ALAN under a wharf in Sydney Harbour (Bolton et al., 2017). The latter showed that, under ALAN, predatory behavior was dramatically greater while abundance was reduced, a similar condition observed during daylight conditions. Interestingly, authors observed a concomitant change in the structure of prey (epifaunal) communities, which suggested an indirect top-down effect of ALAN. It is interesting to note that the presence of night lighting of artificial structures might elicit variable effects on fish depending on the positioning of the light source. Lamps positioned over the structure, for example, create an unnatural high contrast between illuminated areas and darker surroundings, these latter found to attract higher densities of bogues (Boops boops) possibly seeking for a refuge (Georgiadis et al., 2014; Mavraki et al., 2016). Further variability is likely related to species-specific responses, as suggested by the almost lack of effects on juvenile bonefish (Szekeres et al., 2017) or on most fish assemblage inhabiting shallow coastal seagrass beds (Martin et al., 2021).

The mentioned studies suggest that the effect of ALAN on single organisms can easily reverberate on entire communities through trophic and non-trophic interactions. Further, manipulative experiments in the Mediterranean rocky intertidal on biofilm assemblages and their main consumers (i.e., littorinid gastropods) highlighted

quite complex and temporally variable effects. On the short term (~1 month), the increase in consumer pressure by grazers compensated for the positive effect of ALAN (27 lux) on microbial primary producers (MPB), indicating that trophic interactions can provide a stabilizing mechanism against the effects of light pollution on early colonizing autotrophs (Maggi & Benedetti-Cecchi, 2018). On the contrary, on longer terms (~3 months), the presence of ALAN negatively affected the density of grazers. Adding to the positive effect on cyanobacteria, ALAN eventually promoted a shift in the composition of epilithic assemblages, characterized by a higher photoautotrophic diversity at the expense of heterotrophic bacteria (Maggi, Bertocci, & Benedetti-Cecchi, 2020; Maggi, Bongiorni, et al., 2020). Although it is clear how pervasive the role of light pollution on rocky shore intertidal communities is, biological and ecological consequences on different species and communities are still largely unknown.

2.3 | Shallow water coral reef ecosystems

Reef building corals are highly photosensitive, and it is already clear that ALAN is a major emerging sensory pollutant concern for shallow coral reef ecosystems. Yet, ALAN is one of the most understudied threats to corals (Figure 6).

Tropical coral reefs are one of the most biodiverse and productive ecosystems on Earth. Their complex framework offers a unique habitat for thousands of associated species, supporting more species per unit area than any other marine ecosystem. In addition, coral reefs also provide important ecosystem services to millions of people (Hughes et al., 2017, 2018); yet, they have been heavily deteriorated worldwide due to poorly managed anthropogenic activities, habitat loss, and climate change (Hughes et al., 2018).

ALAN has been detected in fringing reefs localized in strongly urbanized locations. Mean night sky brightness levels at reef locations (see Ayalon, Rosenberg, et al., 2021) show that many coral reef areas worldwide are affected by ALAN. An applicable example is the coast in the Gulf of Eilat/Aqaba in the Red Sea, where most of the studies on the effects of ALAN on reef building corals have been conducted (e.g., Ayalon et al., 2019; Levy et al., 2020; Tamir et al., 2017). This region is heavily light polluted, and the light reflected from the cities surrounding the reef can be seen from space (Tamir et al., 2017).

Scleractinian corals in symbiosis with dinoflagellates (Symbiodiniaceae) are the foundation species for the formation of shallow water tropical coral reefs. Because their endosymbiotic partners perform photosynthesis—and up to 95% of the photosynthates can be translocated to the coral host for its metabolic needs—this symbiosis is at the basis of the success of such tropical environments (LaJeunesse et al., 2018; Muscatine et al., 1981; Trembley et al., 2012). The aspects of such symbiosis contribute to a higher susceptibility to ALAN, since corals and their symbionts are

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highly photosensitive and are mostly found in shallow, clear water with relatively high light levels (Rosenberg et al., 2019).

Light is detected by corals through light-sensing molecules, such as cryptochromes (CRY)-proteins that can convert light leading to changes in intracellular levels of important second messengers (Levy et al., 2007). The light/dark cycle regulates many cellular processes, the dark period being crucial for stress recovery and repair, especially for the photosynthetic symbionts (Hill et al., 2011; Levy et al., 2020). Natural periodic illumination (both solar and lunar) is a critical factor in cueing important processes of coral reproduction (Ayalon, Rosenberg, et al., 2021; Lin et al., 2021). In addition, the blue light spectrum—for example, present in LED lights—play a key role in coral growth, symbiont density, Chlorophyll a content, and photosynthetic rates (D'Angelo et al., 2008; Wijgerde et al., 2014). Therefore, it is not surprising that the exposure of corals to ALAN has been shown to have detrimental effects on coral/symbionts metabolism and reproduction (see below). Yet, ALAN is one of the most understudied threats to corals.

Molecular evidence in corals on the effects of ALAN generally match those of more complex organisms, mainly mammalians. Rosenberg et al. (2019) used transcriptomic analysis to compare corals of the species *Acropora eurystoma* growing under natural light cycles and under ALAN (50–40 lux). Authors demonstrated many pathways that were altered, with approximately 25 times more differentially expressed genes that regulate cell cycle, cell proliferation, cell growth, and protein synthesis under ALAN.

Physiological and biochemical investigations on Red Sea corals reported significant deleterious effects of ALAN. Ayalon et al. (2019) first demonstrated that *Acropora eurystoma* and *Pocillopora damicornis* experienced oxidative stress and photosynthetic impairment after exposure to different wavelengths of ALAN (40-35 lux). Subsequently, a more detailed study by Levy et al. (2020) showed that the extent of deleterious effects of ALAN (40-30 lux) on the

symbiotic association (loss of symbionts and Chlorophyll content) and physiology of *Turbinaria reniformis* and *Stylophora pistillata* was aligned with the severity of the oxidative stress condition experienced by the species. The same study also presented preliminary evidence that corals presenting higher basal levels of antioxidant capacity, such as *Turbinaria reniformis*, may be more resistant to ALAN. Ayalon, Benichou, et al. (2021) also reported ALAN can influence Symbiodinaceae cultures and demonstrated different physiological responses according to the algae type. More specifically, *Clodocopium* type showed to be generally more sensitive compared to *Durusdinium* type, presenting decreases in the maximum quantum efficiency of PSII, in the mitotic index, and in total chlorophyll content after exposure to ALAN with illumination level up to 5 lux.

Regarding the effects on reproduction, recent studies reported large effects of ALAN on the gametogenesis and the synchronization of gamete release in prominent coral species from the Indo-Pacific Ocean. The gametogenesis cycle of Acropora millepora and Acropora digitofera was delayed or masked by exposure to ALAN (~15 lux), leading to unsynchronized gamete release (Ayalon, Rosenberg, et al., 2021). Dim light during the night also suppressed spawning in coral Dipsastrea specisosa (Lin et al., 2021). Importantly, this later study showed that the period of darkness between sunset and moonrise is essential to trigger synchronized mass spawning. Additionally, Tamir et al. (2020) reported a 30% decrease in coral settlement success due to ALAN (~20 lux). Such results are alarming for the future of coral reefs. More than 80% of Scleractinian corals are broadcasting spawners (Baird et al., 2009), and asynchronous reproduction caused by ecological speciation could lead to reproductive isolation and prevent gene flow between differential lit coral communities (Rosenberg et al., 2019). Further, successful gamete production and fertilization, development of viable offspring, and survival of coral recruits are undoubtedly some of the most relevant processes for replenishing degraded reefs (Ayalon, Rosenberg, et al., 2021; Harii et al., 2010).

In addition to the symbiosis established with endosymbiotic dinoflagellates, corals are associated with prokaryotic symbionts. In fact, the coral host, and their microbiome (microalgal and prokaryotic symbionts) show a tightly intertwined metabolic activity (Thompson et al., 2015). Coral-associated prokaryotic microbes are taxonomically and functionally diverse and are key for maintaining the health of the holobiont (Hernandez-Agreda et al., 2018; Krediet et al., 2013; Olson et al., 2009). To date, only one study explored the effects of ALAN on the coral microbiome. Baquiran et al. (2020) showed that the overall microbial community structure of the coral Acropora digitifera remained stable under ALAN (~15 lux). However, it is important to note that bacteria that could use light for energy production (chlorophototrophic members of the phylum Proteobacteria), as well as those that are associated with the phototrophic symbionts of the coral, increased in abundance under light pollution conditions. Possibly, the higher abundance of symbiont-associated microbes is linked to greater abundance and activity of the dinoflagellate symbionts under short-term exposure to ALAN (Baquiran et al., 2020).

As for other species inhabiting coral reefs, only effects on fishes have been investigated so far. Fobert et al. (2019, 2020) showed a negative impact of ALAN (~15 lux) on the reproductive success of the common clownfish Amphiprion ocellaris, with an increased interval between spawning events and a smaller size of eggs in comparison to dark conditions; in addition, hatching success was affected both by the presence of ALAN and by its spectral composition, with a more negative effect of cool-white in comparison to warm-white light. After hatching, other life stages might represent critical intervals for the persistence of fish populations under light pollution. O'Connor et al. (2019) highlighted larvae of Acanthurus triostegus living 10 days post-settlement under ALAN conditions (~20-25 lux) experienced higher mortality rates by the end of the experiment, although growing faster and heavier than control ones. Furthermore, a long-term study conducted in the wild (Schligler et al., 2021) showed that environmentally relevant intensity of ALAN (~4.3 lux) is also

able to reduce survival and growth of juveniles of the anemonefish Amphiprion chrysopterus), compared with individuals exposed to natural night illuminance (by moonlight). Finally, a recent study on the blue green Chromis viridis highlighted complex sub-acute effects of ALAN on adult fish, with predator threat able to alter the increased metabolism of both specific tissues and whole organism observed under ALAN conditions (~100 lux). Although an evolving area of study, it is already clear that ALAN is a major emerging sensory pollutant of concern for shallow coral reef ecosystems. ALAN acts as a chronic disturbance, and corals under such pressure may not be able to perform their normal cyclic behaviors (Rosenberg et al., 2019). Therefore, ALAN may impact the future of coral reefs, eventually contributing to global coral reef decline.

2.4 | Pelagic environment organisms

Oceans are vast, three-dimensional, and mostly influenced by currents. Pelagic organisms are not attached to a substrate, hence the direct effects from ALAN in the open oceans are likely to be different from those on a beach or on the seafloor. Still, recent work suggests that lights from ships may have an impact on organisms in both the epipelagic and even mesopelagic zones (Figure 7).

Light influences pelagic organisms in many ways, and artificial light may have a strong impact in their behavior (Blaxter & Currie, 1967). Prey availability, limiting the initiation and magnitude of phytoplankton blooms and mortality through visual predation are some examples of how artificial light may have an impact. In general, most zooplankton are negatively phototactic (Forward Jr, 1988), migrating to depth during daylight to avoid the threat of visual predation, and surfacing at night to feed. This behavior is called Diel Vertical Migration (DVM) (Brierley, 2014). DVM is a characteristic feature of all the world's



FIGURE 7 Lights from a ship working in the dark (Credit; Mike Snyder).

oceans and is considered the largest synchronized movement of biomass on the planet (Hayes, 2003). It is thus an important factor in structuring pelagic communities. At the same time, as this process is mediated by light (Ringelberg, 2010), it may potentially also be strongly affected by artificial light (Berge et al., 2020). At latitudes characterized by midnight sun during summer and polar night during winter, DVM was generally assumed not to occur except during spring and autumn periods when there is a clear day-night cycle (Berge et al., 2009). This view, however, was recently challenged. Instead of an ecosystem that enters a resting state during the polar night, we now recognize a highly active ecosystem characterized by continuous activity and biological interactions across all trophic levels and taxonomic groups (Berge et al., 2015). Importantly, even at the darkest periods of the year, light is still the primary regulative factor for most of these interactions, including vertically migrating zooplankton (Last et al., 2016; Ludvigsen et al., 2018).

It is important to note that behavioral responses to artificial light vary among taxa. While some species are known to be attracted to light, with herring *Clupea harengus* (Stickney, 1969), krill (Krafft & Krag, 2021; Utne-Palm et al., 2018), snow crab *Chionoecetes opilio* (Nguyen et al., 2020), and grey mullet *Mugil cephalus* (Marchesan et al., 2005) as well-known examples, others are known to avoid light. North Atlantic and Arctic copepods (Ludvigsen et al., 2018), Atlantic cod *Gadus morhua* (Utne-Palm et al., 2018), and sea bream *Sparus*

auratus (Marchesan et al., 2005), all species that are commercially important, have been shown to avoid ALAN. A study from the Red Sea, in which a ROV equipped with LED lamps were used to herd mesopelagic scattering layers, similarly concluded that artificial light attracted some species and repelled others (Kaartvedt et al., 2019). Also, recent studies from the high Arctic Archipelago of Svalbard have shown that artificial light from both ships and instrumentation may have a strong impact on organisms down to at least 200 m depth (Berge et al., 2020). The artificial lights in this case were measured to 2.2 µmol photons $m^{-2} s^{-1}$ at the sea surface. However, with field experiments carried out across nearly 8 degrees of latitude, differences in response to artificial light varied both qualitatively and quantitatively in a way that could not be explained by species composition alone. Hence, in addition to interspecific differences in responses to light (Ryer et al., 2009), intraspecific variation could also complicate interpretations of responses to artificial light (Berge et al., 2020).

Effect of light pollution and ALAN in the open ocean is difficult to assess. By default, sampling in the open ocean is often biased, as organisms are not attached or restricted to a specific "site" or physical habitat. And except for acoustic instruments, most sampling technologies include the use of either artificial light or the instrument itself creates a shadow that might influence the organisms (see Box 2). To examine the potential effect of artificial light is, thus, very difficult to do with traditional methodologies that use artificial

BOX 2 How research activities can interfere and create bias due to artificial light

Despite a growing body of literature reporting behavioral disturbance of marine organisms exposed to artificial light, external light sources remain widely used in oceanography and marine ecology studies. Advances in optical technology, combined with the increased desire to use non-lethal observation approaches, have driven the development of new sensors and instruments to document marine ecosystems (Bicknell et al., 2016), but these instruments generally require an external light source. For example, Optical probes such as the Underwater Vision Profiler (Picheral et al., 2010), the Laser-Optical Plankton Counter (Basedow et al., 2013; Herman & Harvey, 2006), the Video Plankton Recorder (Sainmont et al., 2014), and the Light frame On-sight Key species Investigation system (Schmid et al., 2016; Schulz et al., 2010) all use light sources and optical sensors to assess the vertical distribution and abundance of zooplankton. Researchers and the industry alike increasingly use high-definition (HD) video cameras or stereo-cameras mounted on trawls to document the catchability of different species or size classes of fish (Boldt et al., 2018; Underwood et al., 2020; Williams et al., 2010). Such camera systems, when used in dim environments, rely on external light sources to distinguish, and identify marine animals at depth. Although previous studies have raised concerns about the impact of artificial visible light on measurements from optical instruments (Benoit-Bird et al., 2010; Boldt et al., 2018; Doya et al., 2014; Trenkel et al., 2004; Widder et al., 2005), these biases have rarely been quantified (Bicknell et al., 2016). Nonetheless, artificial lighting is assumed to be the main source of biases in fish surveys using cameras and underwater vehicles (Rooper et al., 2015; Ryer et al., 2009; Stoner et al., 2008). The use of red light has been suggested for marine surveys requiring external light sources because it is assumed that most species do not react as much to red light as to shorter wavelengths, such as blue or green (Boldt et al., 2018; Rooper et al., 2015; Widder et al., 2005). In support of this hypothesis, Peña et al. (2020) and Underwood et al. (2020) deployed oceanographic probes equipped with different light colors and showed that mesopelagic (200-1000 m) fish avoid white, blue and green, but not red light. This, however, was recently challenged by Geoffroy et al. (2021) who, using hull-mounted echosounders above an acoustic probe or a baited video camera, each equipped with light sources of different colors (white, blue, and red), demonstrated that pelagic organisms in Arctic and temperate regions strongly avoid artificial light, including visible red light (575-700 nm), from instruments lowered in the water column. Light levels varied within the range of 11-18 μW cm⁻² for the colors tested (see Geoffroy et al., 2021 for details). The density of organisms decreased by up to 99% when exposed to artificial light and the distance of avoidance varied from 23 to 94 m from the light source, depending on colors, irradiance levels and, possibly, species communities (Geoffroy et al., 2021).

light to function. A significant "effect" of ALAN in the open ocean is therefore not restricted to direct effects but will also to a large degree include accuracies and artifacts in the measurements itself.

2.5 | Seabirds

Light pollution causes massive mortality events on seabird fledglings, involving more than 70 species, some of them severely threatened. More subtle effects of ALAN are still poorly understood (Figure 8).

Seabirds are defined as avian species for which a large proportion of the population relies on the marine environment for at least part of the year (Croxall et al., 2012). For example, petrels spend most of their life at sea and only visit land for breeding (Brooke, 2004), while some gull species can spend most of their lives outside the ocean. Within marine biota, seabirds are the only animals mastering the three environments: marine, terrestrial, and aerial. Thus, there is a trade-off among the adaptations of seabirds to cope with different environments. Seabirds are one of the most threatened groups of birds (Dias et al., 2019). According to the International Union for Conservation of Nature (IUCN) Red List criteria, around 31% of all seabird species are globally threatened and 47% have declining population trends.

Environmental conditions, such as light, can rapidly change. Many seabird species nest underground or visit their breeding colonies at night while foraging during daylight. In addition, many seabirds forage by diving in the water column, where the light spectrum changes rapidly with depth (Regular et al., 2011). Seabirds must adapt to rapid changes in intensity and spectra of light to survive. In fact, visual systems of diving seabirds are more sensitive to blue light, the light that penetrates more in-depth (Martin, 2017). The increase of artificial light levels has been identified as a threat for seabirds and, particularly, petrels (Dias et al., 2019; Rodríguez et al., 2019).

Seabirds encounter the most light-polluted areas on land, mostly coastal areas close to or within their nesting colonies. These light sources, such as streetlights, road lights and lighthouses, attract and disorient birds during their flights between colony and foraging sites at sea. Many seabirds visit colonies at night presumably to avoid predation by diurnal predators (Bourgeois et al., 2008; Rubolini et al., 2015). Visit frequency of breeders is influenced by moon cycles probably because of moonlight level variation (Riou & Hamer, 2008; Rodríguez et al., 2016). Thus, breeders could be deterred from visiting colonies when lights are turned on close to their nests or colonies. A recent experimental study on breeders of Manx shearwater Puffinus puffinus demonstrated that adult breeders flying over the colony were deterred by blue and green light in comparison to red light (Syposz et al., 2021). In addition, the number of birds flying over the colony decreased with the duration and intensity of light treatments (Syposz et al., 2021). However, experimental studies on the effect of color and intensity of light (3, 15, and 100 lux) on the colony attendance of the smallest and only penguin whose activity on land is strictly nocturnal, the little penguin Eudyptula minor,



FIGURE 8 Cory's shearwater *Calonectris borealis* fledgling grounded by artificial light in Tenerife, Canary Islands (Credit; Beneharo Rodríguez).

demonstrated that penguins preferred lit paths over dark paths to reach their nests (Rodríguez et al., 2018).

From a conservation point of view and in a short-scale term, the most negative consequence of light pollution is direct mortality caused mainly on fledglings of underground-nesting seabird species (Rodríguez, Holmes, et al., 2017). This phenomenon is known as fallout, and it occurs in all the oceans and seas across the world. Although reasons are unknown (Atchoi et al., 2020), fledglings of many petrel species, but also Alcids and some sea ducks, are attracted and/or disorientated by lights during their maiden flights from their nests toward the ocean (Rodríguez, Holmes, et al., 2017). This leads to grounding and hitting infrastructures, for example, wires, antennas, buildings, or even the ground, causing injuries and fatal victims. If they survive the first collisions, they are vulnerable to other threats, such as vehicle collisions, predation by domestic animals, or traps where they die of inanition or dehydration because they are usually unable to take off. Rescue programs are initiatives aiming to mitigate lightinduced mortality. To reach this, they call for the public implication in rescuing and reporting birds grounded at lit areas, main towns, and cities. Most birds admitted to rescue programs survive (~90%), but the fraction of grounded birds that never are found or reported through

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these collaborative initiatives is unknown. Some studies indicate that around 40% of birds are never rescued (Ainley et al., 2001; Rodríguez et al., 2014). Artificial light could also affect the natural colony attendance of breeders visiting colonies at night.

Increasing ALAN levels in and around nesting colonies could impact seabird's breeding behavior and, consequently, chick provisioning. In an overnight weight gain study, Scopoli's shearwater *Calonectris diomedea* chicks situated closer to a high-light intensity disturbance (i.e., disco event) gained less weight compared to conspecifics from nests further away (Cianchetti-Benedetti et al., 2018). Such effects were not perceivable at fledging, but it is expected that a higher frequency of disturbance events could affect chicks' fitness and, even, breeding success.

Beyond the coastline, seabirds can also encounter extremely light-polluted areas associated to offshore oil and gas platforms, vessels, or light-enhanced fisheries, for example. Our knowledge on light-induced mortality at sea is quite limited (Gjerdrum et al., 2021; Glass & Ryan, 2013; Merkel & Johansen, 2011; Ronconi et al., 2015; Ryan et al., 2021), although we know that adults can also be involved in attraction episodes. Clear monitoring protocols and independent trained observers, who could rely on technological advances, such as telemetry, thermal cameras, acoustic recordings, and radar, are essential to record episodic seabird-light interactions at sea (Ronconi et al., 2015).

The increase of light pollution levels under water (see above) widens the photic zone at night, but also during dawn and dusk. Thus, both at neritic and oceanic waters the increase of light levels by ALAN could enhance the foraging of pursuit-diving visual predator seabirds, such as murres and penguins (Cannell & Cullen, 1998; Elliott & Gaston, 2015; Regular et al., 2011).

Artificial lights can also concentrate prev, which seabirds then take advantage of. Several gull species have been reported to increase their foraging opportunities on marine, coastal and terrestrial lit areas. For example, fishing vessels usually use light to concentrate fish and squid. In the Mediterranean Sea, lights of fishing vessels favor the capture of fish by the Audouin's gull Ichthyaetus audouinii by illuminating the sea surface, concentrating fish, and locating shoals (Arcos & Oro, 2002). Similarly, Brown-hooded gulls Larus maculipennis predate on Isopoda, Polychaeta, fish larvae, and crustaceans, which are concentrated under artificial lights on Argentinian coastal piers (Leopold et al., 2010). On land, Black-backed gull Larus dominicanus can take advantage of Cerambicidae beetles attracted to artificial lights at sawmills (Pugh & Pawson, 2016). Gulls can also prey on seabirds and such predation can be facilitated by artificial light. At Benidorm Island (western Mediterranean), Yellow-legged gulls Larus michahellis increased predation on European stormpetrels Hydrobates pelagicus after light levels increased by a new light installation in the nearby Benidorm city (Oro et al., 2005).

Although there is a certain consensus about the higher pervasiveness of blue light for wildlife and, particularly, seabirds (Longcore et al., 2018; Rodríguez, Dann, & Chiaradia, 2017; Syposz et al., 2021), more research is needed on the spectrum of light in the perception of seabirds. Similarly, reductions in the duration of lights, by means of smart-lighting or part-night lighting, could help to mitigate light

pollution impacts, but current understanding is insufficient to underpin sound recommendations for all species. For example, more research is needed to assess the threshold levels from which a response is triggered as well as the relative intensity with ambient light levels (e.g., during full and new moon nights). The distance from light sources and seabirds at which they are attracted or disorientated must be influenced by light intensity. Thus, determining distances at which individuals are safe is crucial for managing breeding colonies and corridors between colonies and the ocean for inland breeding species. GPS tracking has revealed that most of Cory's shearwater *Calonectris borealis* fledglings are grounded on areas distant <16 km from their nests (Rodríguez et al., 2015, 2022).

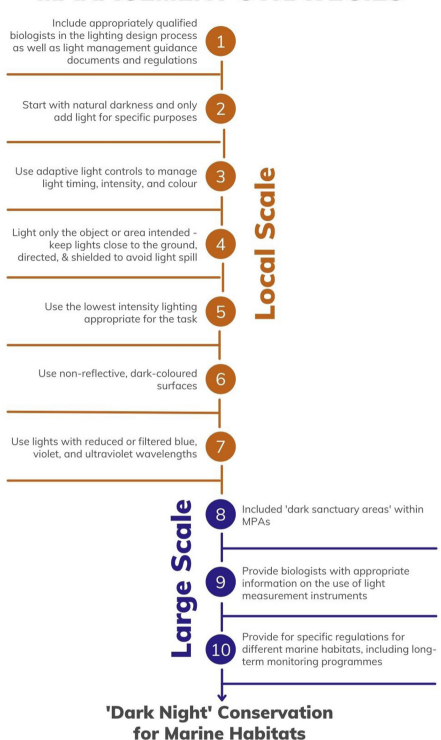
3 | CONSERVATION GUIDELINES AND STRATEGIES

Some few countries today—Spain (Catalonia), Chile, France, and Italy (Piedmont)—are trying to establish laws to regulate light pollution. Most of the applicable documents addressing ALAN are guidelines, Codes or Standards issued by regulators, advisory committees, non-government organizations (NGO) or professional bodies with no legal basis for enforcement of recommendations. Many of the professional body guidance documents are targeted at lighting engineers or designers, provide little detail regarding ALAN management and mitigation for the protection of sensitive receptors and must be purchased at a substantial cost (e.g., AS/NZS 4282, 2019). In addition, regulators, advisory committees, and NGOs typically focus on a single sensitive receptor such as bats, birds or dark sky conservation for astronomy or star gazing (City of Calgary, 2011; NSW, 2016; Voigt et al., 2018).

Following the adoption of the Australian National Light Pollution Guidelines for Wildlife, including Marine Turtles, Seabirds and Migratory Shorebirds (Commonwealth, 2020; CMS, 2020a, 2020b) by the UNEP Convention of Migratory Species in 2020, the CMS Secretariat has expanded on the issue with the release of a follow-up review of ALAN impacts on migratory species not covered by the Australian guidelines (CMS, 2021). This document summarized some of the available international laws and guidelines that address ALAN. Except for the CMS guidelines (CMS, 2020b), laws, standards and codes all relate to terrestrial based ALAN and sensitive receptors. Guidance for addressing the impacts and management of artificial light in the marine environment does not currently exist in the (English) grey literature or publications. Recognizing the difficulties in setting specific assessment trigger values for the broad range of variables influencing the impact of light on wildlife, the CMS guidelines recommend a case-by-case approach to ALAN impact assessment, management, and mitigation. The conservation strategies adopted will vary depending on the sensitive receptor (foraging and migrating birds, hatchling sea turtles migrating offshore, plankton, fish, marine mammals etc.), the sensitivity of the receptor to different wavelengths of light (e.g., whales do not see color), the light sources (ships at anchor offshore, offshore oil and gas facilities and flaring,

MANAGEMENT STRATEGIES

FIGURE 9 Ten golden rules for dark night conservation for marine habitats.



ports and marinas, slow moving dredge vessels etc.), the light features (wavelength/color, intensity, shielding, flaring gas flow rates, elevation) and variables such as turbidity, water depth, clouds, dust, aerosols, moon phase, fog, day length, all acting in combination to influence the visibility of the light. Standard best practice guidelines for outdoor lighting design for the protection of sensitive receptors including wildlife have been published (ADSA, 2021; CMS, 2020b;

IDA, 2021) and include: (i) Start with natural darkness and only add light for specific purposes; (ii) use adaptive light controls to manage light timing, intensity, and color; (iii) light only the object or area intended, (iv) use the lowest intensity lighting appropriate for the task; (v) use non-reflective, dark-colored surfaces; and (vi) use lights with reduced or filtered blue, violet, and ultraviolet wavelengths. Of these, the most important is avoiding short wavelength blue light

due to its ubiquitous visibility across a wide range of taxa, as well as its higher capacity to penetrate the water column, shielding light to prevent light spill into the water or sky, and minimizing light intensity.

A correct application of such rules, for the management of "dark nights" in the marine environment, needs the consultation of appropriately qualified biologists, which should be included in the lighting design process as well as light management guidance documents and regulations. At a larger scale, qualified researchers should help identify appropriate "dark sanctuary areas" within MPAs and providing for specific regulations in different marine habitats, including long-term monitoring programs. Considering the best practices guidelines and their correct application, we propose here "Ten golden rules for dark night conservation for marine habitats" (See Figure 9).

4 | CONCLUSION AND PERSPECTIVES

Progress in our understanding of ALAN impacts in marine ecosystems has accelerated dramatically over the last 5 years. The number of species, habitats and ecological processes with documented responses now present a compelling case for ALAN as a globally widespread pollutant that is reshaping nature across our coastlines. The field remains however, recently emergent, and numerous knowledge gaps exist that if addressed would aid in the prediction and mitigation of ALAN impacts in the sea. Here, we highlight key questions for future research to be addressed:

- 1. What is the impact of artificial light at sea on marine wildlife populations?
- 2. What is the contribution of marine traffic (i.e., mobile light sources from vessels) and traffic management (i.e., fixed light sources from navigation markers) to marine light pollution, and what is their potential to impact marine biodiversity?
- 3. What are the indirect impacts of ALAN in marine ecosystems through species interactions and trophic cascades?
- 4. Are there intergenerational impacts of ALAN?
- 5. What are the best practice techniques to monitor and measure biologically meaningful (i.e., radiometric as opposed to photometric) light, both underwater and on land, at both fine (meters) and broad (kilometers) scales?
- 6. What are the thresholds (intensity, wavelength, exposure time) that elicit biological responses in marine species?
- 7. How does the disruption of moonlight cycles by artificial skyglow impact circalunar rhythms in marine organisms?
- 8. What is the impact of coastal ALAN on marine ecosystem
- Does ALAN impact long distance mass migrations of marine megafauna?
- 10. Is there temporal variability (e.g., monthly, or seasonal) in ALAN impacts on marine ecosystems?
- 11. Are there additive/interactive effects between ALAN and other anthropogenic disturbances?

Addressing these questions will provide insight into the full extent of ALAN impacts in marine ecosystems, their consequences for society, and options for mitigating them.

CONFLICT OF INTEREST

Authors declare that they have no competing interests.

DATA AVAILABILITY STATEMENT

Data sharing is not applicable to this article as no new data were created or analyzed in this study.

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REFERENCES

- ADSA. (2021). Retrieved from https://www.australasiandarkskyallian ce.org/best-practice-lighting
- AS/NZS 4282:2019. (2019). Control of the obtrusive effects of outdoor lighting. *Standards Australia*. Retrieved from https://infostore.saiglobal.com/en-us/standards/as-nzs-4282-2019-1141358_saig_as_as_2703687/
- Ainley, D. G., Podolsky, R., Deforest, L., & Spencer, G. (2001). The status and population trends of the Newell's Shearwater on Kaua'i: Insights from modeling. *Studies in Avian Biology*, *22*, 108–123.
- Arcos, J. M., & Oro, D. (2002). Significance of nocturnal purse seine fisheries for seabirds: A case study off the Ebro Delta (NW Mediterranean). Marine Biology, 141, 277–286.
- Atchoi, E., Mitkus, M., & Rodríguez, A. (2020). Is seabird light-induced mortality explained by the visual system development? Conservation Science and Practice, 2, e195. https://doi.org/10.1111/csp2.195
- Aulsebrook, A. E., Jechow, A., Krop-Benesch, A., Christopher, C. M., Longcore, T., Perkin, E. K., & van Grunsven, R. H. A. (2022). Nocturnal lighting in animal research should be replicable and reflect relevant ecological conditions. *Biology Letter*, 18, 20220035.
- Ayalon, I., Benichou, J. I. C., Avisar, D., & Levy, O. (2021). The endosymbiotic coral algae Symbiodinaceae are sensitive to a sensory pollutant: Artificial Light at Night, ALAN. Frontier in Physiology, 12, 695083.
- Ayalon, I., Marangoni, L. F. B., Benichou, J. I. C., Avisar, D., Levy, O. (2019). Red Sea corals under artificial light pollution at night (ALAN) undergo oxidative stress and photosynthetic impairment. *Global Change Biology*, 25, 4194–4207.
- Ayalon, I., Rosenberg, Y., Benichou, J. I., Campos, C. L. D., Sayco, S. L. G., Nada, M. A. L., Baquiran, J. I. P., Ligson, C. A., Avisar, D., Conaco, C., Kuechly, H. U., Kyba, C. C. M., Cabaitan, P. C., & Levy, O. (2021). Coral gametogenesis collapses under artificial light pollution. Current Biology, 31, 413–419.
- Baird, A. H., Guest, J. R., & Willis, B. L. (2009). Systematic and biogeographical patterns in the reproductive biology of scleractinian corals. Annual Review of Ecology, Evolution, and Systematics, 40, 551–571.
- Baquiran, J. I. P., Nada, M. A. L., Campos, C. L. D., Sayco, S. L. G., Cabaitan, P. C., Rosenberg, Y., Ayalon, I., Levy, O., & Conaco, C. (2020). The

- Prokaryotic Microbiome of Acropora digitifera is stable under short-term artificial light pollution. *Microorganisms*, 8, 1566.
- Barentine, J. C. (2019). Methods for assessment and monitoring of light pollution around ecologically sensitive sites. *Journal of Imaging*, 5, e5050054
- Bascom, W. (1980). Waves and beaches: The dynamics of the ocean surface.

 Doubleday & Co.
- Basedow, S. L., Tande, K. S., Norrbin, M. F., & Kristiansen, S. A. (2013). Capturing quantitative zooplankton information in the sea: Performance test of laser optical plankton counter and video plankton recorder in a Calanus finmarchicus dominated summer situation. *Progress in Oceanography*, 108, 72–80.
- Båtnes, A. S., Miljeteig, C., Berge, J., Greenacre, M., & Johnsen, G. (2013). Quantifying the light sensitivity of Calanus spp. during the polar night: potential for orchestrated migrations conducted by ambient light from the sun, moon, or aurora borealis? *Polar Biology*, 38, 51-65.
- Benoit-Bird, K. J., Moline, M. A., Schofield, O. M., Robbins, I. C., & Waluk, C. M. (2010). Zooplankton avoidance of a profiled open-path fluorometer. *Journal of Plankton Research*, 32, 1413–1419.
- Berge, J., Ambrose, W. G., Jr., Darnis, G., Last, K. S., Leu, E., Cohen, J. H., Johnsen, G., Moline, M. A., Cottier, F., Varpe, Ø., Shunatova, N., Bałazy, P., Morata, N., Massabuau, J.-C., Falk-Petersen, S., Kosobokova, K., Hopp, C. J. M., ... Callesen, T. A. (2009). Diel vertical migration of Arctic zooplankton during the polar night. *Biology Letter*, 5, 69-72.
- Berge, J., Daase, M., Renaud, P. E., Cottier, F., Last, K. S., Varpe, O., Leu, E., Søreide, J., Eiane, K., Falk-Petersen, S., Willis, K., Nygård, H., Vogedes, D., Griffiths, C., Johnsen, G., Lorentzen, D., & Brierley, A. S. (2015). Unexpected levels of biological activity during the polar night offer new perspectives on warming Arctic. *Current Biology*, 25, 2555–2561.
- Berge, J., Geoffroy, M., Daase, M., Cottier, F., Priou, P., Cohen, J. H., Johnsen, G., McKee, D., Kostakis, I., Renaud, P. E., Vogedes, D., Anderson, P., Last, K. S., & Gauthier, S. (2020). Artificial light during the polar night disrupts Arctic fish and zooplankton behaviour down to 200 m depth. Communication Biology, 3, 102.
- Bicknell, A. W. J., Godley, B. J., Sheehan, E. V., Votier, S. C., & Witt, M. J. (2016). Camera technology for monitoring marine biodiversity and human impact. Frontiers in Ecology and the Environment, 14, 424–432
- Blaxter, J. H. S., & Currie, R. I. (1967). The effect of artificial lights on acoustic scattering layers in the ocean. Symposia of the Zoological Society of London, 19, 1–14.
- Boldt, J. L., Williams, K., Rooper, C. N., Towler, R. H., & Gauthier, S. (2018). Development of stereo camera methodologies to improve pelagic fish biomass estimates and inform ecosystem management in marine waters. Fisheries Research, 198, 66–77.
- Bolton, D., Mayer-Pinto, M., Clark, G. F., Dafforn, K. A., Brassil, W. A., Becker, A., & Johnston, E. L. (2017). Coastal urban lighting has ecological consequences for multiple trophic levels under the sea. *Science of the Total Environment*, 576, 1–9.
- Bourgeois, K., Dromzée, S., Vidal, É., & Legrand, J. (2008). Yelkouan shear-water *Puffinus yelkouan* presence and behaviour at colonies: Not only a moonlight question. *Comptes Rendus Biologies*, 331, 88–97.
- Bregazzi, P. K., & Naylor, E. (1972). The locomotor activity rhythm of *Talitrus saltutor* (Montagu) (Crustacea, Amphipoda). *Journal of Experimental Biology, 57*, 375–391.
- Brierley, A. S. (2014). Diel vertical migration. *Current Biology*, 24, R1074-R1076.
- Brooke, M. (2004). Albatrosses and petrels across the World. Oxford University Press.
- Brown, A., & McLachlan, A. (1990). *Ecology of sandy shores*. Elsevier, 328 pp.
- Bruno, J. F., & Bertness, M. D. (2001). Habitat modification and facilitation in benthic marine communities. In M. D. Bertness, S. D. Gaines,

- & M. E. Hay (Eds.), Marine community ecology (pp. 221–251). Sinauer Associates.
- Buckley, L., Caldarone, E., & Ong, T. L. (1999). RNA-DNA ratio and other nucleic acid-based indicators for growth and condition of marine fishes. *Hydrobiologia*, 401, 265–277.
- Cannell, B. L., & Cullen, J. M. (1998). The foraging behavior of Little Penguins Eudyptula minor at different light levels. *Ibis*, 140, 467-471.
- Celano, L., Sullivan, C., Field, A., & Salmon, M. (2018). Seafinding revisited: How hatchling marine turtles respond to natural lighting at a nesting beach. *Journal of Comparative Physiology A*, 204, 1007–1015.
- Chícharo, M. A., & Chícharo, L. (2008). RNA:DNA ratio and other nucleic acid derived indices in marine ecology. *International Journal of Molecular Sciences*, 9, 1453–1471.
- Cianchetti-Benedetti, M., Becciu, P., Massa, B., & Dell'Omo, G. (2018). Conflicts between touristic recreational activities and breeding shearwaters: Short-term effect of artificial light and sound on chick weight. European Journal of Wildlife Research, 64, 19.
- CIE. (2018). ISO23539:2005(E)/CIE S 010/E:2004 (2005). Photometry The CIE System of Physical Photometry. Standard; CIE; Vienna, Austria.
- City of Calgary. (2011). Bird-friendly urban design guidelines. Land use planning and policy. Retrieved from https://www.calgary.ca/pda/pd/current-studies-and-ongoing-activities/urban-design.html
- CMS. (2020a). Resolution 13.5 Light Pollution Guidelines for Wildlife.

 Adopted by the Conference of the Parties at its 13th Meeting (Gandhinagar, February 2020). Retrieved from https://www.cms.int/sites/default/files/document/cms_cop13_res.13.5_light-pollutionguidelines_e.pdf
- CMS. (2020b). Annex to Resolution 13.5 Light Pollution Guidelines National Light Pollution Guidelines for Wildlife including marine turtles, seabirds and migratory shorebirds. Retrieved from https://www.cms.int/sites/default/files/document/cms_cop13_res.13.5_annex_e.pdf
- CMS. (2021). Convention on the Conservation of Migratory Species of Wild Animals 5th Meeting of the Sessional Committee of the CMS Scientific Council (ScC-SC5) Online, 28 June 9 July 2021 UNEP/CMS/ScC-SC5/Inf.7 IMPACT OF LIGHT POLLUTION ON DIFFERENT TAXA OF MIGRATORY SPECIES (Prepared by Ms. Laetitia Nunny on behalf of the CMS Secretariat) 87 pp. Retrieved from https://www.cms.int/sites/default/files/document/cms_scc-sc5_inf.7_impact-of-light-pollution-on-migratory-species_e.pdf
- Colman, L. P., Lara, P. H., Bennie, J., Broderick, A. C., de Freitas, J. R., Marcondes, A., Witt, M. J., & Godley, B. J. (2020). Assessing coastal artificial light and potential exposure of wildlife at a national scale: The case of marine turtles in Brazil. *Biodiversity and Conservation*, 29, 1135–1152.
- Commonwealth of Australia. (2020). National Light Pollution guidelines for Wildlife, including marine turtles, seabirds and migratory shorebirds. 111pp. Retrived from https://www.environment.gov.au/biodiversity/publications/national-light-pollution-guidelines-wildlife
- Croxall, J. P., Butchart, S. H. M., Lascelles, B., Stattersfield, A. J., Sullivan, B., Symes, A., & Taylor, P. (2012). Seabird conservation status, threats and priority actions: A global assessment. *Bird Conservation International*, 22, 1–34.
- Cruz, L. M., Shillinger, G. L., Robinson, N. J., Tomillo, P. S., & Paladino, F. V. (2018). Effect of light intensity and wavelength on the in-water orientation of olive ridley turtle hatchlings. *Journal of Experimental Marine Biology and Ecology*, 505, 52–56.
- D'Angelo, C., Denzel, A., Vogt, A., Matz, M. V., Oswald, F., Salih, A., Nienhaus, G. U., & Wiedenmann, J. (2008). Blue light regulation of host pigment in reef-building corals. *Marine Ecology Progress Series*, 364, 97-106.
- Davies, T. W., & Smyth, T. (2018). Why artificial light at night should be a focus for global change research in the 21st century. *Global Change Biology*, 24, 872–882.

- Davies, T. W., Coleman, M., Griffith, K. M., & Jenkins, S. R. (2015). Nighttime lighting alters the composition of marine epifaunal communities. Biology Letter, 11, 20150080.
- Davies, T. W., Duffy, J. P., Bennie, J., & Gaston, K. J. (2014). The nature, extent, and ecological implications of marine light pollution. Frontiers in Ecology and the Environment, 12, 347–355.
- Davies, T. W., Duffy, J. P., Bennie, J., & Gaston, K. J. (2016). Stemming the tide of light pollution encroaching into marine protected areas. Conservation Letters, 9, 164-171.
- Davies, T. W., McKee, D., Fishwick, J., Tidau, S., & Smyth, T. (2020). Biologically important artificial light at night on the seafloor. Science Report, 10, 12545.
- Dias, M. P., Martin, R., Pearmain, E. J., Burfield, I. J., Small, C., Phillips, R. A., Yates, O., Lascelles, B., Borboroglu, P. G., & Croxalla, J. P. (2019). Threats to seabirds: A global assessment. Biological Conservation, 237, 525-537.
- Dimitriadis, C., Fournari e Konstantinidou, I., Sourbes, L., Koutsoubas, D., & Mazaris, A. D. (2018). Reduction of sea turtle population recruitment caused by nightlight: Evidence from the Mediterranean region. Ocean Coast Management, 153, 108-115.
- Domenici, P., Torres, R., & Manríquez, P. H. (2017). Effects of elevated carbon dioxide and increased temperature on locomotion and the repeatability of lateralization of a keystone marine mollusk. The Journal of Experimental Biology, 220, 667-676.
- Doya, C., Aguzzi, J., Matabos, M., Mihaly, S., & Canals, M. (2014). Diel behavioral rhythms in sablefish (Anoplopoma fimbria) and other benthic species, as recorded by the Deep-sea cabled observatories in Barkley canyon (NEPTUNE-Canada). Journal of Marine Systems, 130, 69-78.
- Duarte, C., Quintanilla-Ahumada, D., Anguita, C., Manríquez, P. H., Widdicombe, S., Pulgar, J., Silva-Rodríguez, E. A., Miranda, C., Manríquez, K., & Quijón, P. A. (2019). Artificial light pollution at night (ALAN) disrupts the distribution and circadian rhythm of a sandy beach isopod. Environmental Pollution, 248, 565-573.
- Elliott, K. H., & Gaston, A. J. (2015). Diel vertical migration of prey and light availability constrain foraging in an Arctic seabird. Marine Biology, 162, 1739-1748.
- Falchi, F., Cinzano, P., Duriscoe, D., Kyba, C., Elvidge, C. D., Baugh, K., Portnov, B. A., Rybnikova, N. A., & Furgoni, R. (2016). The new world atlas of artificial night sky brightness. Science Advances, 2, 1-26.
- Fallaci, M., Aloia, A., Audoglio, M., Colombini, I., Scapini, F., & Chelazzi, L. (1999). Differences in behavioral strategies between two sympatric talitrids (Amphipoda) inhabiting an exposed sandy beach of the French Atlantic Coast. Estuarine, Coastal and Shelf Science, 48,
- Fallaci, M., Aloia, A., Colombini, I., & Chelazzi, L. (2002). Population dynamics and life history of two Phaleria species (Coleoptera, Tenebrionidae) living on the Tyrrhenian sandy coast of central Italy. Acta Oecologica, 23, 69-79.
- Fanini, L., Hughes, L. E., Springthorpe, R., Tosetto, L., & Lowry, J. K. (2016). Surface activity patterns of macrofauna on pocket, tidal beaches: insights into the role of wrack and artificial lighting. Regional Studies in Marine Science, 7, 63-71.
- Fobert, E. K., Burke da Silva, K., & Swearer, S. E. (2019). Artificial light at night causes reproductive failure in clownfish. Biology Letters, 15, 20190272.
- Forward Jr, R. B. J. (1988). Diel vertical migration: zooplankton photobiology and behaviour. Oceanography and Marine Biology: An Annual Review, 26, 361-393.
- Fritsches, K. A. (2012). Australian loggerhead sea turtle hatchlings do not avoid yellow. Marine and Freshwater Behaviour and Physiology, 45. 79-89.
- Gaston, K. J., Ackermann, S., Bennie, J., Cox, D. T., Phillips, B. B., de Miguel, A. S., & Sanders, D. (2021). Pervasiveness of biological impacts of artificial light at night. Integrative & Comparative Biology, 61, 1098-1110.

- Geoffroy, M., Langbehn, T., Priou, P., Varpe, Ø., Johnsen, G., Le Bris, A., Fisher, J. A. D., Daase, M., Cohen, J., & Berge, J. (2021). Pelagic organisms avoid white, blue, and red artificial light from scientific instruments. Scientific Reports, 11, 14941.
- Georgiadis, M., Mavraki, N., Koutsikopoulos, C., & Tzanatos, E. (2014). Spatio-temporal dynamics and management implications of the nightly appearance of Boops boops (Acanthoptervgii, Perciformes) iuvenile shoals in the anthropogenically modified Mediterranean littoral zone. Hydrobiologia, 734, 81-96.
- Gjerdrum, C., Ronconi, R., Turner, K., & Hamer, T. (2021). Bird strandings and bright lights at coastal and offshore industrial sites in Atlantic Canada. Avian Conservation and Ecology, 16, 22.
- Glass, J., & Ryan, P. (2013). Reduced seabird night strikes and mortality in the Tristan rock lobster fishery. African Journal of Mariene Science, 35. 589-592.
- Gonzalez, S. A., Yañez-navea, K., & Muñoz, M. (2014). Effect of coastal urbanization on sandy beach coleoptera Phaleria maculata (Kulzer, 1959) in northern Chile. Marine Pollution Bulletin, 83, 265 - 274
- Hänel, A., Posch, T., Ribas, S. J., Aubé, M., Duriscoe, D., Jechow, A., Kollath, Z., Lolkema, D. E., Moore, C., Schmidt, N., & Spoelstra, H. (2018). Measuring night sky brightness: Methods and challenges. Journal of Quantitative Spectroscopy and Radiative Transfer, 205, 278-290.
- Harii, S., Yamamoto, M., & Hoegh-Guldberg, O. (2010). The relative contribution of dinoflagellate photosynthesis and stored lipids to the survivorship of symbiotic larvae of the reef-building corals. Marine Biology, 157, 1215-1224.
- Hayes, G. C. (2003). A review of the adaptive significance and ecosystem consequences of zooplankton diel vertical migrations. Hydrobiologia, 503, 163-170.
- Hays, G. C., Mackay, A., Adams, C. R., Mortimer, J. A., Speakman, J. R., & Boerema, M. (1995). Nest site selection by sea turtles. Journal of the Marine Biological Association of the United Kingdom, 75, 667-674.
- Helfman, G., Collette, B., Facey, D., & Bowen, B. (2009). The Diversity of Fish: Biology, Evolution and Ecology. Willey-Blackwell, A John Wiley & Sons, Ltd. Publication.
- Herman, A. W., & Harvey, M. (2006). Application of normalized biomass size spectra to laser optical plankton counter net intercomparisons of zooplankton distributions. Journal of Geophysical Research: Oceans, 11, C05S05.
- Hernandez-Agreda, A., Leggat, W., Bongaerts, P., Herrera, C., & Ainsworth, T. D. (2018). Rethinking the coral microbiome: Simplicity exists within a diverse microbial biosphere. MBio, 9, e00812-18.
- Hill, R., Brown, C. M., DeZeeuw, K., Campbell, D. A., & Ralph, P. J. (2011). Increased rate of D1 repair in coral symbionts during bleaching is insufficient to counter accelerated photo-inactivation. Limnology and Oceanography, 56, 139e146.
- Hughes, T. P., Barnes, M. L., Bellwood, D. R., Cinner, J. E., Cumming, G. S., Jackson, J. B., Kleypas, J., van de Leemput, I. A., Lough, J. M., Morrison, T. H., Palumbi, S. R., van Nes, E. H., & Scheffer, M. (2017). Coral reefs in the Anthropocene. Nature, 546, 82-90.
- Hughes, T. P., Anderson, K. D., Connolly, S. R., Heron, S. F., Kerry, J. T., Lough, J. M., Baird, A. H., Baum, J. K., Berumen, M. L., Bridge, T. C., Claar, D. C., Eakin, C. M., Gilmour, J. P., Graham, N. A. J., Harrison, H., Hobbs, J.-P. A., Hoey, A. S., Hoogenboom, M., Lowe, R. J., ... Wilson, S. K. (2018). Spatial and temporal patterns of mass bleaching of corals in the Anthropocene. Science, 359, 80-83.
- IDA. (2021). Retrived from https://www.darksky.org/our-work/lighting/ lighting-principles/
- Jaramillo, E., Contreras, H., Duarte, C., & Avellanal, M. H. (2003). Locomotor activity and zonation of upper shore arthropods in a sandy beach of north central Chile. Estuarine, Coastal and Shelf Science, 58, 177-197.
- Jaramillo, E., Dugan, J. E., Hubbard, D. M., Manzano, M., & Duarte, C. (2021). Ranking the ecological effects of coastal armoring on

3652486, 2022, 18, Downloaded from https://onlinelibrary.wiley.com/doi/10.1111/gcb.16264 by Readcube (Labtiva Inc.), Wiley Online Library on [29/10/2022]. See the Terms and Conditions

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and-conditions) on Wiley Online Library for rules of use; OA articles are governed by the applicable Creative Commons I

- mobile invertebrates across intertidal zones on sandy beaches. Science of the Total Environment, 755, 142573.
- Jechow, A., & Hölker, F. (2019). How dark is a river? Artificial light at night in aquatic systems and the need for comprehensive night-time light measurements. Wiley Interdisciplinary Reviews: Water, 6, e1388.
- Jechow, A., Kyba, C., & Hölker, F. (2019), Beyond all-sky: Assessing ecological light pollution using multi-spectral full-sphere fisheve lens imaging. Journal of Imaging, 5, 46.
- Jenkins, S. R., & Hartnoll, R. G. (2001). Food supply, grazing activity and growth rate in the limpet Patella vulgata L.: A comparison between exposed and sheltered shores. Journal of Experimental Marine Biology and Ecology, 258, 123-139.
- Kaartvedt, S., Røstad, A., Opdal, A. F., & Aksnes, D. L. (2019). Herding mesopelagic fish by light. Marine Ecology Progress Series, 625, 225-231.
- Kamrowski, R. L., Limpus, C., Jones, R., Anderson, S., & Hamann, M. (2014). Temporal changes in artificial light exposure of marine turtle nesting areas. Global Change Biology, 20, 2437-2449.
- Kamrowski, R. L., Limpus, C., Moloney, J., & Hamann, M. (2012). Coastal light pollution and marine turtles: Assessing the magnitude of the problem. Endangered Species Research, 19, 85-98.
- Kamrowski, R. L., Limpus, C., Pendoley, K., & Hamann, M. (2015). Influence of industrial light pollution on the sea-finding behavior of flatback turtle hatchlings. Wildlife Research, 41, 421-434.
- Kelly, I., Leon, J. X., Gilby, B. L., Olds, A. D., & Schlacher, T. A. (2017). Marine turtles are not fussy nesters: A novel test of small-scale nest site selection using structure from motion beach terrain information. PeerJ, 5, e2770.
- Kennedy, F., Naylor, E., & Jaramillo, E. (2000). Ontogenetic differences in the circadian locomotor activity rhythm of the talitrid amphipod crustacean Orchestoidea tuberculata. Marine Biology, 137, 511-517.
- Keough, M. J., & Raimondi, P. T. (1995). Responses of settling invertebrate larvae to bioorganic films: Effects of different types of films. Journal of Experimental Marine Biology and Ecology, 185, 235–253.
- King, P., & Symes, D. (2004). Potential loss of GNP and GSP from a failure to maintain California's beaches. Shore and Beach, 72, 3-8.
- Krafft, B. A., & Krag, L. A. (2021). Antarctic krill (Euphausia superba) exhibit positive phototaxis to white LED light. Polar Biology, 44, 483-489.
- Krediet, C. J., Ritchie, K. B., Paul, V. J., & Teplitski, M. (2013). Coralassociated micro-organisms and their roles in promoting coral health and thwarting diseases. Proceedings of the Royal Society B: Biological Sciences, 280, 20122328.
- LaJeunesse, T. C., Parkinson, J. E., Gabrielson, P. W., Jeong, H. J., Reimer, J. D., Voolstra, C. R., & Santos, S. R. (2018). Systematic revision of Symbiodiniaceae highlights the antiquity and diversity of coral endosymbionts. Current Biology, 28, 2570e2580.
- Last, K. S., Hobbs, L., Berge, J., Brierley, A. S., & Cottier, F. (2016). Moonlight drives ocean-scale mass vertical migration of zooplankton during the Arctic winter. Current Biology, 26, 244-251.
- Leopold, M. F., Philippart, C. J. M., & Yorio, P. (2010). Nocturnal feeding under artificial light conditions by Brown-hooded Gull (Larus maculipennis) in Puerto Madryn harbour (Chubut Province, Argentina). Hornero, 25, 55-60.
- Levy, O., Appelbaum, L., Leggat, W., Gothlif, Y., Hayward, D. C., Miller, D. J., & Hoegh-Guldberg, O. (2007). Light-responsive cryptochromes from a simple multicellular animal, the coral Acropora millepora. Science, 318, 467-470.
- Levy, O., Marangoni, L. F. B., Benichou, J. I., Rottier, C., Béraud, E., Grover, R., & Ferrier-Pagès, C. (2020). Artificial light at night (ALAN) alters the physiology and biochemistry of symbiotic reef building corals. Environmental Pollution, 266, 114987.
- Limpus, C., & Kamrowski, R. L. (2013). Ocean-finding in marine turtles: The importance of low horizon elevation as an orientation cue. Behaviour, 15, 863-893.
- Lin, C-H, Takahashi, S., Mulla, A. J., Nozawa, Y. (2021). Moonrise timing is key for synchronized spawning in coral Dipsastraea speciosa. PNAS, 118, e2101985118.

- Lohmann, K. J., Witherington, B. E., Lohmann, C. M., & Salmon, M. (2017). Orientation, navigation, and natal beach homing in sea turtles. In The biology of sea turtles (pp. 108-135). CRC Press.
- Longcore, T., & Rich, C. (2004). Ecological light pollution. Frontiers in Ecology and the Environment, 2, 191-198.
- Longcore, T., Duriscoe, D., Aubé, M., Jechow, A., Kyba, C., & Pendoley, K. L. (2020). Commentary: Brightness of the night sky affects loggerhead (Caretta caretta) sea turtle hatchling misorientation but not nest site selection. Frontiers in Marine Science, 7, 706. https://doi. org/10.3389/fmars.2020.00706
- Longcore, T., Rodríguez, A., Witherington, B., Penniman, J. F., Herf, L., & Herf, M. (2018). Rapid assessment of lamp spectrum to quantify ecological effects of light at night. Journal of Experimental Zoology Part A: Ecological Genetics and Physiology, 329, 511-521.
- Luarte, T., Bonta, C. C., Silva-Rodriguez, E. A., Quijón, P. A., Miranda, C., Farias, A. A., & Duarte, C. (2016). Light pollution reduces activity, food consumption and growth rates in a sandy beach invertebrate. Environmental Pollution, 218, 1147-1153.
- Lucas, M., Salmon, M., Fritz, E., & Wyneken, J. (1992). Sea finding by hatchling sea turtles: Role of brightness, silhouette and beach slope as orientation cues. Behaviour, 122, 56-77.
- Ludvigsen, M., Berge, J., Geoffroy, M., Cohen, J. H., De La Torre, P. R., Nornes, S. M., Singh, H., Sørensen, A. J., Daase, M., & Johnsen, G. (2018). Use of an autonomous surface vehicle reveals small-scale diel vertical migrations of zooplankton and susceptibility to light pollution under low solar irradiance. Science Advances, 4, eaap9887.
- Lynn, K. D., Flynn, P. T., Manríquez, K., Manríquez, P. H., Pulgar, J., Duarte, C., & Quijon, P. A. (2021). Artificial light at night alters the settlement of acorn barnacles on a man-made habitat in Atlantic Canada. Marine Pollution Bulletin, 163, 111928.
- Lynn, K. D., Quintanilla-Ahumada, D., Anguita, C., Widdicombe, S., Pulgar, J., Manríquez, P. H., Quijón, P. A., & Duarte, C. (2021). Disruption and recovery: Artificial Light at Night (ALAN) alter the activity and feeding behavior of sandy beach Amphipods from Atlantic Canada. Science of the Total Environment, 780, 146568.
- Maggi, E., & Benedetti-Cecchi, L. (2018). Trophic compensation stabilizes marine primary producers exposed to artificial light at night. Marine Ecology Progress Series, 606, 1-5.
- Maggi, E., Bongiorni, L., Fontanini, D., Capocchi, A., Dal Bello, M., Giacomelli, A., & Benedetti-Cecchi, L. (2020). Artificial light at night erases positive interactions across trophic levels. Functional Ecology, 34, 694-706.
- Maggi, E., Bertocci, I., & Benedetti-Cecchi, L. (2020). Light pollution enhances temporal variability of photosynthetic activity in mature and developing biofilm. Hydrobiologia, 847, 1793-1802.
- Mavraki, N., Georgiadis, M., Koutsikopoulos, C., & Tzanatos, E. (2016). Unravelling the nocturnal appearance of bogue Boops boops shoals in the anthropogenically modified shallow littoral. Journal of Fish Biology, 88, 2060-2066.
- McLachlan, A., Defeo, O., Jaramillo, E., & Short, A. D. (2013). Sandy beach conservation and recreation: guidelines for optimizing management strategies for multi-purpose use. Ocean and Coastal Management, 71, 256-268.
- Manríquez, K., Quijon, P. A., Manríquez, P. H., Miranda, C., Pulgar, J., Quintanilla- Ahumada, D., & Duarte, C. (2021). Artificial Light at Night (ALAN) negatively affects the settlement success of two prominent intertidal barnacles. Marine Pollution Bulletin, 168, 112416.
- Manríquez, P. H., Jara, M. J., Gonzalez, G. P., Seguel, M., Quijon, P. A., Widdicombe, S., Pulgar, J. M., Quintanilla-Ahumada, D., Anguita, C., & Duarte, C. (2021). Effects of artificial light at night and predator cues on foraging and predator avoidance in the keystone inshore mollusc Concholepas concholepas. Environmental Pollution, 280, 116895.
- Manríquez, P. H., Jara, M. E., Díaz, M. I., Quijon, P. A., Widdicombe, S., Manríquez, K., Quintanilla-Ahumada, D., & Duarte, C. (2019).

- Artificial light pollution influences behavioral and physiological traits in a keystone predator species, *Concholepas concholepas*. *Science of the Total Environment*, 661, 543–552.
- Manríquez, P. H., Jara, M. E., Mardones, M. L., Torres, R., Navarro, J. M., Lardies, M. A., Vargas, C. A., Duarte, C., & Lagos, N. A. (2014). Ocean acidification affects predator avoidance behaviour but not prey detection in the early ontogeny of a keystone species. *Marine Ecology Progress Series*, 502, 157–167.
- Manríquez, P. H., Lagos, N. A., Jara, M. E., & Castilla, J. (2009). Adaptive shell color plasticity during the early ontogeny of an intertidal keystone predator. *Proceedings of the National Academy of Sciences of the United States of America*, 106, 16298–16303.
- Marchesan, M., Spoto, M., Verginella, L., & Ferreiro, E. A. (2005).Behavioral effects of artificial light on fish species of commercial interest. Fisheries Research, 73, 171–185.
- Martin, G. R. (2017). The sensory ecology of birds. Oxford University Press.
- Martin, C. W., Reynolds, L. K., Scheffel, W. A., Tiffany, S., & Kopetman, S. (2021). Diel variability and influence of artificial light on fish and macroinvertebrate communities in gulf of mexico seagrass beds. Estuaries and Coasts, 44, 431–441.
- Menge, B. A., & Branch, G. M. (2001). Rocky intertidal communities. In M. D. Bertness, S. D. Gaines, & M. E. Hay (Eds.), Marine community ecology (pp. 221–251). Sinauer Associates.
- Merkel, F. R., & Johansen, K. L. (2011). Light-induced bird strikes on vessels in Southwest Greenland. Marine Pollution Bulletin, 62, 2330–2336.
- Meschini, E., Gagliardo, A., & Papi, F. (2008). Lunar orientation in sandhoppers is affected by shifting both the moon phase and the daily clock. *Animal Behaviour*, 76, 25–35.
- Mundy, C. N., & Babcock, R. C. (1998). Role of light intensity and spectral quality in coral settlement: implications for depth-dependent settlement? *Journal of Experimental Marine Biology and Ecology*, 223, 235–255.
- Muscatine, L., Grossman, D., & Doino, J. (1981). Release of symbiotic algae by tropical sea anemones and corals after cold shock. Marine Ecology Progress Series, 77, 233e243.
- Nagarkar, S., Williams, G. S., Subramanian, G., & Saha, S. K. (2004). Cyanobacteria-dominated biofilms: A high quality food resource for intertidal grazers. *Hydrobiologia*, 519, 89–94.
- Navarro-Barranco, C., & Hughes, L. E. (2015). Effects of light pollution on the emergent fauna of shallow marine ecosystems: Amphipods as a case study. *Marine Pollution Bulletin*, 94, 235–240.
- Newell, G. E. (1965). The eye in Littorina littorea. *Proceedings of the Zoological Society of London*, 144, 75–86.
- Nguyen, T. T., Le, M.-H., Doan, N. X., Pham, H. Q., Vu, M. T. T., & Dinh, K. V. (2020). Artificial light pollution increases the sensitivity of tropical zooplankton to extreme warming. *Environmental Technology and Innovation*, 20, 101179.
- NSW. (2016). NSW Planning and Environment. Dark Sky Planning Guideline Protecting the observing conditions at Siding Spring. Department of planning and Environment. Retrieved from https://www.planning.nsw.gov.au/-/media/Files/DPE/Guidelines/dark-sky-planning-guideline-2016-06.pdf
- O'Connor, J. J., Fobert, E. K., Besson, M., Jacob, H., & Lecchini, D. (2019). Live fast, die young: Behavioural and physiological impacts of light pollution on a marine fish during larval recruitment. *Marine Pollution Bulletin*, 146, 908–914.
- Olson, N., Ainsworth, T., Gates, R., & Takabayashi, M. (2009). Diazotrophic bacteria associated with Hawaiian Montipora corals: Diversity and abundance in correlation with symbiotic dinoflagellates. *Journal of Experimental Marine Biology and Ecology*, 371, 140–146.
- Oro, D., de León, A., Minguez, E., & Furness, R. W. (2005). Estimating predation on breeding European storm-petrels (*Hydrobates pelagicus*) by yellow-legged gulls (*Larus michahellis*). *Journal of Zoology*, 265, 421–429.

- Paine, R. T. (1966). Food web complexity and species diversity. American Naturalist, 100, 65.
- Patrício, A. R., Varela, M. R., Barbosa, C., Broderick, A. C., Airaud, M. B. F., Godley, B. J., & Catry, P. (2018). Nest site selection repeatability of green turtles, *Chelonia mydas*, and consequences for offspring. *Animal Behaviour*, 139, 91–102.
- Peña, M., Cabrera-Gámez, J., & Domínguez-Brito, A. C. (2020). Multifrequency and light-avoiding characteristics of deep acoustic layers in the North Atlantic. Marine Environmental Research. 154, 104842.
- Pendleton, L., Kildow, J., & Rote, J. W. (2006). The non-market value of beach recreation in California. Shore and Beach. 74, 34–37.
- Pendoley, K., & Kamrowski, R. L. (2015). Influence of horizon elevation on the sea-finding behaviour of hatchling flatback turtles exposed to artificial light glow. Marine Ecology Progress Series, 529, 279-288.
- Picheral, M., Guidi, L., Stemmann, L., Karl, D. M., Iddaoud, G., & Gorsky, G. (2010). The underwater vision profiler 5: An advanced instrument for high spatial resolution studies of particle size spectra and zooplankton. *Limnology and Oceanography-Methods*, 8, 462–547.
- Pugh, A. R., & Pawson, S. M. (2016). Artificial light at night potentially alters feeding behaviour of the native southern black-backed gull (*Larus dominicanus*). Notornis, 63, 37–39.
- Pulgar, J., Zeballos, D., Vargas, J., Aldana, M., Manriquez, P. H., Manriquez, K., Quijón, P. A., Widdicombe, S., Anguita, C., Quintanilla, D., & Duarte, C. (2019). Endogenous cycles, activity patterns and energy expenditure of an intertidal fish is modified by artificial light pollution at night (ALAN). Environmental Pollution, 244, 361–366.
- Qian, P. Y., Lau, S. C. K., Dahms, H. U., Dobretsov, S., & Harder, T. (2007). Marine biofilms as mediators of colonization by marine macroorganisms: Implications for antifouling and aquaculture. *Marine Biotechnology*, 9, 399–410.
- Quintanilla-Ahumada, D., Quijón, P. A., Pulgar, J., Manríquez, P. H., García-Huidobro, M. R., & Duarte, C. (2021). Exposure to artificial light at night (ALAN) alters RNA: DNA ratios in a sandy beach coleopteran insect. *Marine Pollution Bulletin*, 165, 112132.
- Quintanilla-Ahumada, D., Quijón, P. A., Manríquez, P. H., Pulgar, J., García-Huidobro, M. R., Miranda, C., Molina, A., Zuloaga, R., & Duarte, C. (2022). Artificial light at night (ALAN) causes variable dose-responses in a sandy beach isopod. *Environmental Science* and Pollution Research, 29, 35977–35985. https://doi.org/10.1007/ s11356-021-17344-2
- Regular, P. M., Hedd, A., & Montevecchi, W. A. (2011). Fishing in the dark: A pursuit-diving seabird modifies foraging behaviour in response to nocturnal light levels. *PLoS One*, 6, e26763.
- Ringelberg, J. (2010). Diel vertical migration of zooplankton in lakes and oceans: Causal explanations and adaptations. Springer Science.
- Riou, S., & Hamer, K. C. (2008). Predation risk and reproductive effort: Impacts of moonlight on food provisioning and chick growth in Manx shearwaters. *Animal Behaviour*, 76, 1743–1748.
- Rivas, M. L., Santidrián Tomillo, P., Uribeondo, J. D., & Marco, A. (2015). Leatherback hatchling sea-finding in response to artificial lighting: interaction between wavelength and moonlight. *Journal of Experimental Marine Biology and Ecology*, 463, 143–149.
- Rodríguez, A., Arcos, J. M., Bretagnolle, V., Dias, M. P., Holmes, N. D., Louzao, M., Provencher, J., Raine, A. F., Ramírez, F., Rodríguez, B., Ronconi, R. A., Taylor, R. S., Bonnaud, E., Borrelle, S. B., Cortés, V., Descamps, S., Friesen, V. L., Genovart, M., Hedd, A., ... Chairadia, A. (2019). Future directions in conservation research on petrels and shearwaters. Frontiers in Marine Science, 6, 94.
- Rodríguez, A., Burgan, G., Dann, P., Jessop, R., Negro, J. J., & Chiaradia, A. (2014). Fatal attraction of short-tailed shearwaters to artificial lights Schierwater, B. (ed.). PLoS One, 9, e110114.
- Rodríguez, A., Chiaradia, A., Wasiak, P., Renwick, L., & Dann, P. (2016). Waddling on the dark side: Ambient light affects attendance behavior of little penguins. *Journal of Biological Rhythms*, 31, 194-204.

- Rodríguez, A., Dann, P., & Chiaradia, A. (2017). Reducing light-induced mortality of seabirds: High pressure sodium lights decrease the fatal attraction of shearwaters. *Journal for Nature Conservation*, 39, 68–72.
- Rodríguez, A., Holmberg, R., Dann, P., & Chiaradia, A. (2018). Penguin colony attendance under artificial lights for ecotourism. *Journal of Experimental Zoology Part A: Ecological and Integrative Physiology*, 329, 457–464.
- Rodríguez, A., Holmes, N. D., Ryan, P. G., Wilson, K.-J., Faulquier, L., Murillo, Y., Raine, A. F., Penniman, J. F., Neves, V., Rodríguez, B., Negro, J. J., Chiaradia, A., Dann, P., Anderson, T., Metzger, B., Shirai, M., Deppe, L., Whe, J., ... Le Corre, M. (2017). Seabird mortality induced by land-based artificial lights. *Conservation Biology*, 31, 986–1001.
- Rodríguez, A., Rodríguez, B., & Negro, J. J. (2015). GPS tracking for mapping seabird mortality induced by light pollution. *Scientific Reports*, 5, 10670.
- Rodríguez, A., Rodríguez, B., Acosta, Y., & Negro, J. J. (2022). Tracking flights to investigate seabird mortality induced by artificial lights. Frontiers in Ecology and Evolution, 9, 786557.
- Ronconi, R. A., Allard, K. A., & Taylor, P. D. (2015). Bird interactions with offshore oil and gas platforms: Review of impacts and monitoring techniques. *Journal of Environmental Management*, 147, 34–45.
- Rooper, C. N., Williams, K., De Robertis, A., & Tuttle, V. (2015). Effect of underwater lighting on observations of density and behavior of rockfish during camera surveys. Fisheries Research, 172, 157–167.
- Rosenberg, Y., Doniger, T., & Levy, O. (2019). Sustainability of coral reefs are affected by ecological light pollution in the Gulf of Aqaba/Eilat. *Communications Biology*, 2, 289.
- Rubolini, D., Maggini, I., Ambrosini, R., Imperio, S., Paiva, V. H., Gaibani, G., Saino, N., & Cecere, J. G. (2015). The effect of moonlight on Scopoli's shearwater *Calonectris diomedea* colony attendance patterns and nocturnal foraging: A test of the foraging efficiency hypothesis. *Ethology*, 121, 284–299.
- Ryan, P. G., Ryan, E. M., & Glass, J. P. (2021). Dazzled by the light: The impact of light pollution from ships on seabirds at Tristan da Cunha. *Ostrich*, *92*, 218–224.
- Ryer, C. H., Stoner, A. W., Iseri, P. J., & Spencer, M. L. (2009). Effects of simulated underwater vehicle lighting on fish behavior. *Marine Ecology Progress Series*, 391, 97–106.
- Sainmont, J., Gislason, A., Heuschele, J., Webster, C. N., Sylvander, P., Wang, M., & Varpe, Ø. (2014). Inter- and intra-specific diurnal habitat selection of zooplankton during the spring bloom observed by Video Plankton Recorder. *Marine Biology*, 161, 1931–1941.
- Salmon, M., & Wyneken, J. (1987). Orientation and swimming behavior of hatchling loggerhead turtles Caretta caretta L. during their offshore migration. Journal of Experimental Marine Biology and Ecology, 109, 137–153.
- Salmon, M. (2003). Artificial night lighting and sea turtles. *Biologist*, 50, 163–168.
- Salmon, M., Tolbert, M. G., Painter, D. P., Goff, M., & Reiners, R. (1995). Behavior of loggerhead sea turtles on an urban beach. II. Hatchling orientation. *Journal of Herpetology*, 29, 568–576.
- Sanders, D., Frago, E., Kehoe, R., Patterson, C., & Gaston, K. J. (2021). A meta-analysis of biological impacts of artificial light at night. *Nature Ecology & Evolution*, 5, 74–81.
- Schlacher, T. A., Lucrezi, S., Connolly, R. M., Peterson, C. H., Gilby, B. L., Maslo, B., Olds, A. D., Walker, S. J., Leon, J. X., Huijbers, C. M., Weston, M. A., Turra, A., Hyndes, G. A., Holt, R. A., & Schoeman, D. S. (2016). Human threats to sandy beaches: A meta-analysis of ghost crabs illustrates global anthropogenic impacts. *Estuarine, Coastal and Shelf Science*, 169, 56-73.
- Schligler, J., Cortese, D., Beldade, R., Swearer, S. E., & Mills, S. C. (2021). Long-term exposure to artificial light at night in the wild decreases survival and growth of a coral reef fish. *Proceedings of the Royal Society B*, 288, 20210454.

- Schmid, M. S., Aubry, C., Grigor, J., & Fortier, L. (2016). The LOKI underwater imaging system and an automatic identification model for the detection of zooplankton taxa in the Arctic Ocean. *Methods in Oceanography*, 15-16, 129-160.
- Schulz, J., Barz, K., Ayon, P., Luedtke, A., Zielinski, O., Mengedoht, D., & Hirche, H. J. (2010). Imaging of plankton specimens with the light frame on-sight key species investigation (LOKI) system. *Journal of the European Optical Society*, 5, 10017s.
- Shimada, T., Meekan, M. G., Baldwin, R., Al-Suwailem, A. M., Clarke, C., Santillan, A. S., & Duarte, C. M. (2021). Distribution and temporal trends in the abundance of nesting sea turtles in the Red Sea. *Biological Conservation*, 261, 109235.
- Smyth, T. J., Wright, A. E., McKee, D., Tidau, S., Tamir, R., Dubinsky, Z., & Davies, T. W. (2021). A global atlas of artificial light at night under the sea. *Elementa: Science of the Anthropocene*, *9*, 00049. https://doi.org/10.1525/elementa.2021.00049
- Stickney, A. P. (1969). Factors influencing the attraction of Atlantic Herring. Fishery Bulletin, 68, 73–85.
- Stoner, A. W., Ryer, C. H., Parker, S. J., Auster, P. J., & Wakefield, W. W. (2008). Evaluating the role of fish behavior in surveys conducted with underwater vehicles. *Canadian Journal of Fisheries and Aquatic Sciences*, 65, 1230–1243.
- Syposz, M., Padget, O., Willis, J., Van Doren, B. M., Gillies, N., Fayet, A. L., Wood, M. J., Alejo, A., & Guilford, T. (2021). Avoidance of different durations, colors, and intensities of artificial light by adult seabirds. *Scientific Reports*, 11, 18941.
- Szekeres, P., Wilson, A. D. M., Haak, C. R., Danylchuk, A. J., Brownscombe, J. W., Elvidge, C. K., Shultz, A. D., Birnie-Gauvin, K., & Cooke, S. J. (2017). Does coastal light pollution alter the nocturnal behavior and blood physiology of juvenile bonefish (Albula vulpes)? *Bulletin of Marine Science*, 93, 491–505.
- Tamir, R., Eyal, G., Cohen, I., & Loya, Y. (2020). Effects of light pollution on the early life stages of the most abundant Northern Red Sea Coral. *Microorganisms*, 8, 193.
- Tamir, R., Lerner, A., Haspel, C., Dubinsky, Z., & Iluz, D. (2017). The spectral and spatial distribution of light pollution in the waters of the northern Gulf of Aqaba (Eilat). Scientific Reports, 7, 42329.
- Thompson, J. R., Rivera, H. E., Closek, C. J., & Medina, M. (2015). Microbes in the coral holobiont: Partners through evolution, development, and ecological interactions. Frontiers in Cellular and Infection Microbiology, 4, 176.
- Thorson, G. (1964). Light as an ecological factor in the dispersal and settlement of larvae of marine bottom invertebrates. *Ophelia*, 1, 167–208.
- Torres, D., Tidau, S., Jenkins, S., & Davies, T. (2020). Artificial skyglow disrupts celestial migration at night. *Current Biology*, 30, R696-R697.
- Trembley, P., Grover, R., Maguer, J.-F., Legendre, L., & Ferrier-Page, C. (2012). Autotrophic carbon budget in coral tissue: A new ¹³C-based model of photosynthates translocation. *The Journal of Experimental Biology*, 215, 1384e1393.
- Trenkel, V. M., Lorance, P., & Mahévas, S. (2004). Do visual transects provide true population density estimates for deepwater fish? *ICES Journal of Marine Science*, *61*, 1050–1056.
- Truscott, Z., Booth, D. T., & Limpus, C. J. (2017). The effect of on-shore light pollution on sea-turtle hatchlings commencing their off-shore swim. *Wildlife Research*, 44, 127–134.
- Ugolini, A., Fantini, T., & Innocenti, R. (2003). Orientation at night: An innate moon compass in sandhoppers (Amphipoda: Talitridae). Proceedings of the Royal Society of London, 270, 279–281.
- Underwood, C. N., Davies, T. W., & Queirós, A. M. (2017). Artificial light at night alters trophic interactions of intertidal invertebrates. *The Journal of Animal Ecology*, 86, 781–789.
- Underwood, M. J., Utne Palm, A. C., Øvredal, J. T., & Bjordal, Å. (2020). The response of mesopelagic organisms to artificial lights. Aquaculture and Fisheries, 6, 519–529.

- Utne-Palm, A. C., Breen, M., Løkkeborg, S., & Humborstad, O. B. (2018).

 Behavioral responses of krill and cod to artificial light in laboratory experiments. *PLoS ONE*, *13*, e0190918.
- Voigt, C. C., Currie, S. E., Fritze, M., Roeleke, M., & Lindecke, O. (2018). Conservation strategies for bats flying at high altitudes. *BioScience*, 68, 427–435.
- Wells, R. (1980). Activity pattern as a mechanism of predator avoidance in two species of acmaeid limpet. *Journal of Experimental Marine Biology and Ecology*, 48, 151–158.
- Widder, E. A., Robison, B. H., Reisenbichler, K. R., & Haddock, S. H. D. (2005). Using red light for in situ observations of deep-sea fishes. Deep-Sea Research Part I, 52, 2077–2085.
- Wijgerde, T., van Melis, A., Silva, C. I. F., Leal, M. C., Vogels, L., Mutter, C., & Osinga, R. (2014). Red light represses the photophysiology of the scleractinian coral Stylophora pistillata. PLoS ONE, 9, e92781.
- Williams, K., Rooper, C. N., & Towler, R. (2010). Use of stereo camera systems for assessment of rockfish abundance in untrawlable areas and for recording pollock behaviour during midwater trawls. *Fishery Bulletin*, 108, 352–362.
- Wilson, P., Thums, M., Pattiaratchi, C., Meekan, M., Pendoley, K., Fisher, R., & Whiting, S. (2018). Artificial light disrupts the nearshore dispersal of neonate flatback turtles Natator depressus. *Marine Ecology Progress Series*, 600, 179–192.
- Wilson, P., Thums, M., Pattiaratchi, C., Whiting, S., Meekan, M., & Pendoley, K. (2021). Nearshore wave characteristics as cues for swimming orientation in flatback turtle hatchlings.

- Journal of Experimental Marine Biology and Ecology, 535, 151475.
- Windle, A. E., Hooley, D. S., & Johnston, D. W. (2018). Robotic vehicles enable high-resolution light pollution sampling of sea turtle nesting beaches. *Frontiers in Marine Science*, *5*, 493.
- Witherington, B. E., & Bjorndal, K. A. (1991a). Influences of artificial lighting on the seaward orientation of hatchling loggerhead turtles *Caretta caretta*. *Biological Conservation*, 55, 139–149.
- Witherington, B. E., & Bjorndal, K. A. (1991b). Influences of wavelength and intensity on hatchling sea turtle phototaxis: Implications for sea-finding behavior. *Copeia*, 1991, 1060–1069.
- Wood, D. W., & Bjorndal, K. A. (2000). Relation of temperature, moisture, salinity, and slope to nest site selection in loggerhead sea turtles. *Copeia*, 2000, 119.
- Wyneken, J., & Salmon, M. (1992). Frenzy and post renzy swimming activity in loggerhead, green, and leatherback hatchling sea turtles. *Copeia*, 2, 478–484.

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