What is the ‘spectral diet’ of humans?
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Our visual perception of the world — seeing form and colour or navigating the environment — depends on the interaction of light and matter in the environment. Light also has a more fundamental role in regulating rhythms in physiology and behaviour, as well as in the acute secretion of hormones such as melatonin and changes in alertness, where light exposure at short-time, medium-time and long-time scales has different effects on these visual and non-visual functions. Yet patterns of light exposure in the real world are inherently messy: we move in and out of buildings and are therefore exposed to mixtures of artificial and natural light, and the physical makeup of our environment can also drastically alter the spectral composition and spatial distribution of the emitted light. In spatial vision, the examination of natural image statistics has proven to be an important driver in research. Here, we expand this concept to the spectral domain and develop the concept of the ‘spectral diet’ of humans.

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Human innovation has produced dramatic changes in our light environment due to artificial lighting. Artificial light comes in many variants: incandescent, fluorescent, and most recently, light emitting diode (LED) lighting. These differ in the spectrum of the emitted light. Except in windowless rooms or rooms with full shading from outside illumination, indoor illumination always reflects a mixture of artificial and natural illumination as shown in Figure 1a.

Light emitted from any source also interacts with various types of matter before reaching an observer. Whether due to Rayleigh scattering in the atmosphere, or specular reflections from the glazed window panels of office buildings (see the example spectrum S3,2 in Figure 1c), light received at the eye is a product of interactions in our environment, which change the originally emitted source spectrum. Material properties alter the light environment in rather complex ways because surfaces in the real-world exhibit complex properties. We do not live in a Lambertian world, in which surfaces are matte. For this reason, estimating the effect of known illuminants on a human observer is non-trivial.

The visual world fortunately exhibits statistical regularities which naturally limit the set of unique combinations of illuminant spectra and reflective surfaces [7,8]. There are other regularities: the sky is in general brighter than the ground, and there is a bias towards dark contrasts in the sky but more balanced contrasts on the ground [9,10]. The same holds true of the spectra we are exposed to and a statistical approach can help to reduce the space of all possible spectra without needing to simulate complex scenes in a physically realistic way. The concept of...
The modern light environment is complex, consisting of spectral data from a unique combination of illuminant spectra and reflected surfaces over space. In this office setting (a), there is both direct and indirect (reflected) light. There are three direct sources: the sky (daylight), the computer screen, and the overhead fluorescent lighting. All other observed spectra are generated by source spectra interacting with objects in the scene, whose reflectance properties are not ideally diffuse (Lambertian). To simulate different representative spectral contributions in the scene (b and c), we assumed reflectance values from the IES TM-30-15 Advanced Calculation Tool [5] and a dataset of 401 illuminant spectra [6]. The resulting spectrum over the pixel area comprises a weighted average of the incident spectra. As the pixel area approaches 1, each pixel corresponds to a unique spectrum in the scene.

standard illuminants (such as D65 ‘noon daylight’ or F11 fluorescent lighting) could be extended to whole spectral scenes, using a discrete basis set of spectral mixtures whose profiles would have been empirically derived and which we would be exposed to at a very high frequency in real life (e.g. see Figure 3b).

What are the visual and non-visual effects of light on humans?
The human visual system comprises a variety of neural mechanisms which allows us to see space, detail, colour, and motion in the world, allows us to make value judgements about objects and actions, and to navigate from one place to another. In Figure 2a, we see how light information from a scene is projected onto the retina and subsequently processed to form an output signal leading to various physiological and psychological effects. Vision and visual perception are adaptive, responding to short-term, mid-term and long-term changes in the visual environment. Exposure to bright light can have long-lasting effects on visual perception. For example, exposure to one hour of sunlight can lead to shifts in colour matches, which can last for as long as five hours [11]. Exposure to filtered lights can also lead to long-lasting changes in colour perception [12]. In the laboratory, exposure to a background light can be used to selectively adapt and suppress specific photoreceptor mechanisms [13].

By contrast, ‘non-visual’ or ‘non-image-forming’ effects are effects of light that are not related to vision or visual perception, such as circadian phase shifting [14], the acute suppression of melatonin [15], the regulation of pupil size [16] and the modulation of alertness [17**]. The term ‘non-visual’, however, is a misnomer, used as a catch-all term to describe a range of biological responses which relate to processes facilitated by the singular visual system, with all light-mediated signals originating from the retina. Similarly, the term ‘non-image-forming’ implies that there is no spatial specificity to these responses, even though from first principles, any photoreceptive system, no matter how broadly tuned it is in space, has a finite receptive field. There is also evidence suggesting that the recently discovered intrinsically
photosensitive retinal ganglion cells (ipRGCs or pRGCs) modulate outputs from rods and cones and even play a role in pattern vision [18,19].

Retinal photoreception proceeds from the absorption of photons by the photoreceptors, initiating a response in the form of an neutral pulse sent via the optic nerve to the suprachiasmatic nucleus (SCN) via a complex pathway where the circadian pacemaker is synchronised by the properties of the light stimulus [20]. The timing of circadian rhythms, regulated by the SCN, is then responsible for regulating sleep onset together with homeostatic processes. Destabilising the circadian pacemaker can result in the disruption of some or all of these processes, which in turn can result in changes in health, alertness, and wellbeing [21].

Retinal photoreceptors can be classified into three broad groups: rods, cones further classified as short (S), medium (M) or long (L) based upon their spectral sensitivity, and photosensitive retinal ganglion cells expressing the photopigment melanopsin. Rods and...
cones contribute directly to the image forming process and indirectly with pRGCs to communicate information about light/dark transitions and light levels [22,23]. A complete understanding of the relative photoreceptor contributions (Figure 2c) and their combination (e.g. through opponency [24]) in the non-image-forming process is still lacking. Models fit to existing datasets [25] indicate that there may be a time-dependent relationship between the relative contributions of L and M cones and melatonin when it comes to the sensitivity of melatonin suppression over time [26]. In addition, there is evidence in primates that S cones may inhibit pRGC excitation [27].

Ocular filters in front of the retina alter the spectrum of light reaching the retina [28], with the lens density being age-dependent [29] and the macular pigment being prominent only in the centre of the retina [30]. Pupil size varies depending on light level, relative photoreceptor activation (e.g. inhibition by S and M cones [31,32]) and many other factors [33] between 2 and 8 mm in diameter (corresponding to a difference in retinal illuminance of only a factor of ~16). As Barlow noted, it can “best be regarded as an inefficient homeostatic device if all it accomplishes is a reduction of input range from 10^11 to 10^10 mW [34]. But one order of magnitude change may make a difference in melatonin suppression [35], which exhibits a melatonin suppression [35], which exhibits a relatively sharp threshold [15]. This characteristic ‘step’ behavior is also reflected in recorded alertness data [36]. Retinal illuminance may also be modulated by squinting and the specific facial features, such as the brow, nose, and eyelid position, of an individual [37]. Finally, the retinal image is displaced frequently by head, eye and trunk movements [38–40].

Why do we care about patterns of light in the environment?

Human exposure to light has changed profoundly. In the US in the 1800s, it is estimated that ~90% of the population worked outside. By 2001, Americans spent 87% of their wake hours indoors [41]. The consequences of this substantial shift are poorly appreciated and the consequences on human health and performance remain unclear. Planners and designers of the built environment alone cannot predict how aspects of the built environment will impact life, but the light environment as a whole is fundamentally related to behavioural, psychophysical, and physiological responses. Light can be measured relatively easily, and new wearable sensors are making large-scale studies more feasible. At the city scale, access to light can be traced back to specific design interventions and strategies, e.g. in streetlighting. Understanding how features of the built environment facilitate light exposure is critical in understanding how modern living could be better aligned with human biology.

The exact impact of light on the totality of visual and non-visual responses is very difficult to predict. For example, a certain light exposure may directly affect physiology and behaviour by leading to an increase in alertness. That same light, however, may shift the circadian phase, which in turn indirectly could modulate alertness over a longer time scale, making an individual more prone to fatigue. Further, there might be an indirect effect of increased fatigue, causing a decrease in subjective wellbeing, all while conditions for visual comfort must be maintained. This ‘nesting’ of different neurophysiological processes and their entangled outcomes illustrates the complexity of separating indirect effects from direct effects. For this reason, it is a key challenge to develop methods of assessing the stability of the circadian system under different illumination patterns over extended periods of time (i.e. days and weeks).

Drawing an analogy to food, we refer to these patterns of light exposure as the ‘spectral diet’. The absolute spectral diet is the pattern of absolute spectral irradiance a human observer might receive over the course of a specific amount of time. The relative spectral diet refers to the spectral composition, or spectral quality, of the light received. Both absolute and relative spectral diets are important. The analogy to food diet here is the following: one might eat, over the course of the waking day, a fixed number of calories, for example, 2000 kcal. This number itself does not tell us how these 2000 kcal were spread across the different macronutrients. Similarly, an illuminance of 2000 lx received during daytime hours does not tell us how the different photoreceptors were affected by it. The concept of the spectral diet can be conceptualised and visualised by a continuum of hyperspectral images stacked together seen over time as illustrated in Figure 2b, which results in spectral data over the image span as visualised in Figure 3a.

We note that interrogating statistical regularities in the environment, and specifically in natural images, has a long history in neuroscience [42–45]. The spectral diet is not simply a characterisation of the spectral environment for its own sake but could potentially also lead to inferences about the properties that an ‘ideal’ – or statistically optimal – observer’s photoreceptive system should have (e.g. spectral tuning, or temporal filtering properties).

How can we measure patterns of light exposure in the environment?

Apart from being a useful concept to think about light input, how can we measure and characterise the spectral diet? Diurnal patterns of light exposure have been measured for healthy adults [46–50] and in psychiatric and neurological diseases and disorders (e.g. Seasonal Affective Disorder [51]; Alzheimer’s Disease [52]) using illuminance sensors. These sensors weigh the spectrum by the photopic luminosity function, which as a combination
of L and M cones does not contain all information required to estimate the non-visual impact of a light.

A recent international standard (CIE S 026/E:2018) describes a retinally referenced framework for quantifying the effects of light from a given radiation or irradiance spectrum [53**]. This standard, based on previous proposals [54], allows for the determination of the extent to which the L, M and S cones, the rods, and melanopsin are activated by a given spectrum. For these calculations, the spectrum needs to be known or alternatively, sensors or imaging systems would need to incorporate the spectral sensitivities described by the standard. Another option is that spectral information could be computationally recovered (to some imperfect precision and accuracy) from a different set of sensors [55–58], or different sensors could be used to recover information about cone, rod and melanopsin excitation [59] without a hyperspectral image.

What are the options of measuring spectral light exposure? Most spectrometers rely on a scanning linear CCD which limits their size and typically affordability. However, new technology using array-based sensors are now able to measure at 5 nm resolution within the visible range. These can be integrated into portable spectrometers worn at the face plane and may provide a more
accurate assessment of the light reaching the eye (though with the caveats mentioned above, such as pupil size). These miniature spectrometers at the time of writing this review (mid 2019) on the market have footprints under 25 mm². Compact circuit boards can be integrated with other sensors measuring activity, temperature, and GPS location to record near-continuous data with a long battery life. Some sample data collected with such a sensor are shown in Figure 3a.

The combination of new in-field calibration methods and weather sealing allow integrated spectral sensors to be used effectively under naturalistic, real-world conditions. The flexibility and relative low-cost of wearable sensors enable researchers to effectively measure light exposure on the neighbourhood and city scales and characterise the spectral diet in a large group of individuals. Data from such measuring campaigns could be then subjected to mathematical and computational models of the circadian system to determine circadian parameters such as amplitude and phase (e.g. [60**]), and be made available to other researchers using data sharing platforms (e.g. FigShare).

**Conclusion**

The relationship between light and human biology is complex. Light intensity, duration, wavelength and timing, along with the individual history of light exposure and the age of the individual all need to be taken into consideration when characterising these relationships and an appropriate ‘spectral diet’. Since the beginning of human evolution, the circadian pacemaker has relied on daylight as the primary environmental zeitgeber. As the human environment changes to accommodate a 24/7 life style, our activity schedules are no longer tied to the availability of daylight. Artificial light, on the other hand, may manipulate timing and periodicity of our internal pacemaker. The statistical regularities in an individual’s spectral diet could be a first step in identifying how the spectral environment determines non-visual responses. But spectral data alone are not enough and for this reason we must also rely on ancillary data such as activity, core body temperature, sleep schedule, melatonin suppression, and others in order to contextualise incident light spectra and constrain models of visual and non-visual function.

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