Divergent photic thresholds in the non-image-forming visual system: entrainment, masking and pupillary light reflex

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Light is the principal cue that entrains the circadian timing system, but the threshold of entrainment and the relative contributions of the retinal photoreceptors—rods, cones and intrinsically photosensitive retinal ganglion cells—are not known. We measured thresholds of entrainment of wheel-running rhythms at three wavelengths, and compared these to thresholds of two other non-image-forming visual system functions: masking and the pupillary light reflex (PLR). At the entrainment threshold, the relative spectral sensitivity and absolute photon flux suggest that this threshold is determined by rods. Dim light that entrained mice failed to elicit either masking or PLR; in general, circadian entrainment is more sensitive by 1–2 log units than other measures of the non-image-forming visual system. Importantly, the results indicate that dim light can entrain circadian rhythms even when it fails to produce more easily measurable acute responses to light such as phase shifting and melatonin suppression. Photosensitivity to one response, therefore, cannot be generalized to other non-image-forming functions. These results also impact practical problems in selecting appropriate lighting in laboratory animal husbandry.

Keywords: circadian; suprachiasmatic nucleus; retina; photoreceptor; wavelength

1. INTRODUCTION

The circadian system plays a critical role in regulating rhythms in physiology and behaviour, and in keeping them synchronized to 24-h rhythms in the environment. The light–dark cycle is the principal cue that entrains the circadian clock in the suprachiasmatic nucleus (SCN) of the hypothalamus. While threshold and relative spectral sensitivity to light have been studied for some indices of circadian function (e.g. phase shifts, melatonin suppression: [1–3]), these have not been described for entrainment. Whether threshold determinations based on responses to acute light pulses extrapolate to entrainment is not known. As a consequence, the photoreceptors that mediate entrainment, and the limits of sensitivity to light of this fundamental circadian function remain unknown.

The circadian clock in the SCN is one component of the broader non-image-forming visual system that also includes control of pupil size and the acute modulation of activity by light [4]. Mammalian photoreceptors include rods, cones and melanopsin-containing intrinsically photosensitive retinal ganglion cells (ipRGC) [5]. The relative contributions of the three photoreceptors to a given physiological or behavioural response depend on wavelength, intensity and duration of light.

The intensity in the diurnal portion of the light:dark cycle (LD) affects how animals entrain. The dimmer the light, the earlier locomotor activity and melatonin secretion begin [6,7]. Dim light in the nocturnal portion of the LD cycle also affects phase angle of entrainment and the rate of resynchronization in jetlag models [8]. Furthermore, some mice switch from nocturnal to diurnal behaviour in dim light [9]. Despite these effects of dim light, the few studies that have addressed entrainment thresholds have employed white light and have studied retinally degenerate and mutant mice, thus precluding both the description of spectral sensitivity of entrainment, and the responsible photoreceptors, in animals with normal retinae [10–15].

In this study, we assessed wheel-running rhythms in mice in diminishing irradiance steps in three wavelengths to identify the irradiance threshold and the photoreceptors that mediate entrainment. These thresholds were compared with those for masking, the pupillary light reflex (PLR), as well as threshold values in the literature for these and other measures of the non-image-forming visual system. The results have practical significance for the types of light used in the laboratory animal husbandry. More importantly, the results show that dim light can entrain circadian rhythms even though it may be too dim to elicit other circadian responses.

2. METHODS

(a) Animals and housing

Male C57Bl/6 mice (Charles River Laboratories, Wilmington, MA, USA) were housed individually in clear polycarbonate...
cages (32 × 14 × 13 cm) on pine shavings, with food (LabDiet 5001, PMI Nutrition, Brentwood, MO, USA) and water available ad libitum. The light cycle was 12 h light and 12 h dark (12 L: 12 D) except during masking when animals were housed in 3.5 L: 3.5 D (DT17C, Intermatic, Spring Grove, IL or XT Table Top Timer, ChronTel, San Diego, CA, USA). Each cage was equipped with a running wheel (11 cm diameter). Environmental noise was masked by white noise (76 dB SPL).

(b) Light
Light-emitting diodes (LEDs) were mounted in an array on the ceiling of each shelf to evenly illuminate the cages, 15 cm below, with blue, green or red light. The inter-bulb distance was 8–9 cm for blue and green LEDs and 4–5 cm for red LEDs. Effective wavelengths (λe) were obtained from the product of the LED emission spectra and each photoreceptor’s sensitivity at each wavelength according to Gavrovskii et al. [16] and are used throughout (blue: λpeak ± spectral half width (mean λe): 464 ± 12 (465 nm); green: 524 ± 23 (518 nm); red: λpeak = 639 ± 9 (635 nm); details in the electronic supplementary material, table S1).

For testing PLR, the eye was illuminated with a single LED (blue: 470 ± 11 (470 nm); green: 520 ± 18 (517 nm); red: 632 ± 10 (626 nm); electronic supplementary material, table S1). Irradiance and photopic illuminance were measured 4 cm from the cage floor (IL1700, International Light Technologies, Peabody, MA, USA). Irradiance was converted and is reported as log photon flux (log(photons cm⁻² s⁻¹)) unless otherwise noted. One group of mice was tested for entrainment thresholds in red and blue light, PLR in red and blue light and masking in blue light. A second group of mice was tested for entrainment in green light, PLR in green light and masking in green and red light.

(c) Responses measured
To measure entrainment, wheel-running activity was monitored remotely by VITALVIEW (Minimitter, Bend, OR, USA), with counts collected in 10 min bins, and plotted in actograms. Daily onset of activity bouts was calculated by CLOCKLAB (Actimetrics, Wilmette, IL, USA), and adjusted actograms. Half-maximal response irradiances (I₅₀) were determined from the best-fit four parameter logistic function (see the electronic supplementary material). The data were fit to Vitamin-A1 spectral sensitivity functions [16] by the method of least squares. A Monte-Carlo simulation was used to generate confidence intervals for the best-fit photopigment templates (see the electronic supplementary material). Differences in response to wavelength were assessed with ANOVA, and pairwise differences assessed with the post hoc Tukey test. First responses to light were assessed by one-sample tests for deviation from 50 per cent (masking), and by one-tailed paired t-tests to compare initial and final pupil size (PLR). Individual differences were assessed by Pearson correlation. These tests were all performed using STATVIEW (SAS Institute, Inc., Cary, NC, USA). Significance was set at the 0.05 level.

3. RESULTS
(a) Entrainment
At each phase shift and step down of irradiance, we evaluated whether mice free-ran or entrained to the new photocycle. Entrainment is manifest here as a transient interval with period less than 24 h followed by a period of 24 h and phase angle of entrainment of 0.5–3 h. This can be seen in the individual activity records and in the plots of mean phase angle relative to lights-off (figure 1a–c). At irradiances that entrain all mice, mean phase angle gets earlier during the re-entrainment interval and stabilizes at a constant phase angle (e.g. green 11.7, figure 1b). At irradiances that entrain only a subset of mice, the contributions of mice that re-entrain and those that continue to free-run are observed in the change in slope of onset time over days (e.g. during intervals at 9.8 (465 nm) and 9.4 (518 nm), figure 1a,b). The per cent of animals entrained at each step is shown in figure 1d.

Table 1 shows the mean minimum light intensities to which mice entrained. Sensitivity to green and blue light was similar, with both thresholds significantly lower than for red (ANOVA, F₂,₉₉ = 52.3, p < 0.001; Tukey test, p < 0.001). Similar results were obtained when the I₅₀ of entrainment was estimated from the per cent of mice entrained at each irradiance step (electronic supplementary material, figure S1). The mean thresholds were used in the studies of masking and PLR described next.

(b) Masking
Masking is a graded response: bright light suppresses wheel running, while dim light has little effect (figure 2). Negative masking is more pronounced at short wavelengths, and positive masking is more pronounced at longer wavelengths (figure 2b). I₅₀ of masking (log photon flux) was 12.4 ± 0.2 for blue, 13.4 ± 0.3 for green and 16.0 ± 0.4 for red (figure 3a; logistic function coefficients a = min = 0%, δ = max = 50%, c = slope = −0.37). There was a significant effect of photon
Figure 1. Representative double-plotted activity records (left panels) and phase angle (group data, right panels) in blue (a) (465 nm, n = 11), green (b) (518 nm, n = 15), and red light (c) (635 nm, n = 16). The log photon flux at each intensity step is marked to the left. In the left panels, lights-off is in grey and the LD transitions marked by lines. Wheel running is indicated by black hatch marks; consecutive days proceed from top to bottom. Onsets on the right side of the actograms are connected for ease of visualization. Last intensity to which masking is never observed at 465 nm; negative masking is observed in dimmer light, especially during the first 30 min of light exposure (518 nm at 9 and 10.4; 635 nm at 11.7 and 12.6). Positive masking is never observed at 465 nm; negative masking is never observed at 635 nm.

Table 1. Irradiance and photopic illuminance at entrainment thresholds.

<table>
<thead>
<tr>
<th>λ (nm)</th>
<th>log photon flux (log photons cm⁻² s⁻¹)</th>
<th>irradiance (nW cm⁻² s⁻¹)</th>
<th>illuminance (lux)</th>
</tr>
</thead>
<tbody>
<tr>
<td>465</td>
<td>9.46 ± 0.19</td>
<td>1.24</td>
<td>0.001</td>
</tr>
<tr>
<td>518</td>
<td>9.13 ± 0.20</td>
<td>0.52</td>
<td>0.0005</td>
</tr>
<tr>
<td>635</td>
<td>11.87 ± 0.19</td>
<td>228</td>
<td>0.4</td>
</tr>
</tbody>
</table>

flux on the degree of masking in both blue (repeated measures ANOVA, \( F_{0.78} = 27.8, p < 0.001 \)) and green light (\( F_{5,50} = 11.8, p < 0.001 \)). At the threshold of entrainment, there was no significant negative masking by light of any wavelength (one-sample test, \( p > 0.05 \)).

In a few mice, negative masking could be observed at irradiances too dim to cause entrainment (mouse in 465 and 635 nm, figure 1).

(c) PLR
Light at the threshold irradiance for entrainment failed to constrict the pupil (figure 3a, one-sample test, \( p > 0.05 \)). Significant constriction as assessed by t-test...
was detected first at log photon fluxes of 10.4 (470 nm),
11.4 (517 nm, but note that \( p = 0.08 \) at 10.4) and 12.4
(626 nm). PLR is a graded response over approximately
five orders of magnitude (figure 3b; repeated measures
ANOVA, 470 nm: \( F_{7,40} = 138; \) 517 nm: \( F_{7,28} = 29; \)
626 nm: \( F_{6,78} = 86; \) all \( p < 0.001 \). \( I_{50/PLR} \) (log photon
flux) was 13.2 ± 0.1 for blue, 13.1 ± 0.1 for green and
15.0 ± 0.1 for red \( (a = 5\% \), \( b = 100\% \), \( d = -0.59 \)).

(d) Spectral sensitivity of non-image-forming
visual functions
Mean entrainment thresholds, \( I_{50/masking} \) and \( I_{50/PLR} \) were
plotted against Govardovskii Vitamin-A1 templates
for the murine rod \( (\lambda_{\text{max}} = 498 \text{ nm}) \), mid-wavelength
cone \( (508 \text{ nm}) \) and ipRGC \( (480 \text{ nm}) \) figure 4a–c). The
ultraviolet cone is minimally sensitive to these wavelengths and is not included [22]. Raw
entrainment threshold data were best fit by an A1 template with a peak at 503 nm. A Monte-Carlo simulation
based on the means and standard deviation for entrainment, or the \( I_{50} \) and standard deviation for masking and
PLR, were used to generate 95% confidence intervals
(CI) for the best-fit A1 templates. With 1000 iterations,
the simulation produced distributions showing the likelihood of finding a best-fit template with given \( \lambda_{\text{max}} \)
(figure 4d). The simulation yielded a mean \( \lambda_{\text{max}} \) of
503 nm for entrainment (CI, 492–514 nm), 482 nm for
masking (CI, 460–504 nm) and 514 nm for PLR
(CI, 510–518 nm).

(e) Comparison of sensitivity thresholds
To understand relative sensitivity among the three
measures, the irradiance–response curves calculated
above were re-plotted by wavelength (figure 5a–c). As measured at both 50 per cent and 10 per cent maximal
response, entrainment is more sensitive than masking
and PLR (figure 5d). A survey of the literature on photosensitivity indicates that the entrainment \( I_{50} \) is 1–2 log
units lower than half-maximal values for other measures of
the non-image-forming visual system in general and
of the circadian system in particular (electronic supplementary material, table S2).

(f) Individual differences
We reasoned that if sensitivity to light is determined by
retinal factors alone, then individuals most sensitive in
one measure should be most sensitive in the others.

Nevertheless, we found no significant correlations
between thresholds for entrainment, masking and PLR
(blue entrainment: versus masking, \( R = -0.41 \), or versus
PLR, \( R = -0.10 \); green entrainment: versus masking,
demonstrates that PLR is mediated primarily by entrainment, much more is known about photoreceptors define the lower limit of entrainment.

With a 95% CI that overlapped with entrainment in our mice was best fit by an A1 template both rods and cones, the relative spectral sensitivity of retinae, clarify the roles of rods and cones. Consistent with these outer layer photoreceptors, but used white light and did not separate the role of rods and cones ([10,12]; but see [11]). The distinct roles of rods and cones are unclear. Two studies point to rods as mediating entrainment in dim light: mice whose photoreception is limited to rods remain capable of entraining to dim light [14,15]. These two studies also showed that the lengthening of free-running period in constant light, another measure of circadian photoreception, does not require cones or melanopsin. By contrast, mice that have rods but lack the mid-wavelength cone do not entrain to LD cycles at 1 lux while controls do entrain, implicating the cone as the limiting photoreceptor [13]. These studies all employed mutant mice in addressing entrainment sensitivity, which may partly explain the discrepancies.

The present data, obtained in wild-type mice with intact retinae, clarify the roles of rods and cones. Consistent with both rods and cones, the relative spectral sensitivity of entrainment in our mice was best fit by an A1 template with a 95% CI that overlapped with $\lambda_{\text{max}}$ for both rods and cones. Thus, spectral data do not differentiate between these photoreceptors. The photon flux at threshold, however, suggests a rod-mediated mechanism. At the threshold of entrainment in 518 nm light, the estimated incident photon flux on the retina is $5E + 8$ photons cm$^{-2}$ s$^{-1}$ (corneal photon flux of $1.4E + 9$ photons cm$^{-2}$ s$^{-1}$, dark adapted pupil size of 4.9 mm$^2$; retina size of 14.5 mm$^2$; [23]). This falls in the dynamic range of rods [24], and is too dim to excite cones [25] or ipRGCs [26]. These data, therefore, strongly suggest that rods define the lower limit of entrainment.

Compared with the photoreceptors mediating entrainment, much more is known about photoreceptors mediating PLR. Work in mutant and wild-type mice demonstrates that PLR is mediated primarily by cones [15,18]. The present results confirm that PLR is cone-mediated.

Negative masking is mediated by ipRGC photoreception, whereas rods and possibly cones, contribute to both positive and negative masking depending on light intensity [17,27,28]. Our masking data support this (figure 2b): negative masking is most pronounced and positive masking is undetectable in blue light that maximally excites ipRGCs, whereas negative masking is absent and positive masking is greatest in red light that has a greater relative effect on rods and cones compared with ipRGCs. The $\lambda_{\text{max}}$ confidence interval for the negative masking spectral sensitivity function was wide and overlapped with both ipRGCs and rods. This is consistent with the strong negative masking in bright light at shorter wavelengths, and also with the low amplitude and transient masking in light too dim to excite ipRGCs. Also, despite large differences in $I_{50}$, there is some overlap in the dynamic ranges between masking and entrainment (figures 1 and 5).

These data have important practical and basic biological implications. On the practical side, they help define appropriate background lighting in laboratory animal husbandry. Dim red light (typically reported as less than 1 lux or less than 0.1 lux) is routinely used to aid behavioural observations and animal care, and constant dim red light is often substituted for constant darkness in circadian studies for the same reason. The justifications include the relatively low sensitivity of rhodopsin to red light, and the inability of dim red light to phase-shift circadian rhythms [2,29]. Nevertheless, the present report shows that dim red light on the order of 0.1–1 lux is an effective entraining signal. Dim red light and constant darkness are not equivalent (e.g. [30]).

With regard to basic biology, the results indicate a range of photic sensitivities across different measures and have implications for our understanding of photoreception. Entrainment is more sensitive to light than are either masking or PLR. The entrainment $I_{50}$ is also lower than half-maximal irradiances reported in the literature for other indices of circadian function, including phase shifting and melatonin suppression (electronic supplementary material, table S2). Even though a given light may be too dim to elicit an acute response from the circadian system, it may nevertheless entrain the circadian system. This is important for understanding the non-image-forming visual system, as phase shifting and melatonin suppression are often used as indices for light’s effect on the circadian system, in both human and animal studies [31,32].

Dim light presented for long durations and detected by rods can entrain circadian rhythms in normal mice with intact retinae. This confirms the sufficiency of rods as mediators of entrainment in dim light [14,15]. The striking difference in photosensitivity among physiological measures shows that photosensitivity of a given measure cannot be easily inferred from surrogate indices. The actions of long-duration dim light are not well understood beyond the behavioural level. Determining the mechanisms by which light detected by rods and cones is integrated remains to be explored.

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REFERENCES


