Ocular and Systemic Diurnal Rhythms in Emmetropic and Myopic Adults

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Purpose. To investigate ocular and systemic diurnal rhythms in emmetropic and myopic adults and examine relationships with light exposure.

Methods. Adult subjects (n = 42, 22–41 years) underwent measurements every 4 hours for 24 hours, including blood pressure, heart rate, body temperature, intraocular pressure (IOP), ocular biometry, and optical coherence tomography imaging. Mean ocular perfusion pressure (MOPP) was calculated. Saliva was collected for melatonin and cortisol analysis. Acrophase and amplitude for each parameter were compared between refractive error groups. Subjects wore a light, sleep, and activity monitor for 1 week before measurements.

Results. All parameters exhibited significant diurnal rhythm (ANOVA, P < 0.05 for all). Choroidal length peaked at 2.42 hours, with a diurnal variation of 12.96 ± 3.71 μm. Axial length peaked at 12.96 hours, with a variation of 25.8 ± 6.6 μm. Melatonin peaked at 3.19 hours during the dark period, while cortisol peaked after light onset at 8.86 hours. IOP peaked at 11.24 hours, with a variation of 4.92 ± 1.57 mm Hg, in antiphase with MOPP, which peaked at 22.02 hours. Amplitudes of daily variations were not correlated with light exposure, and rhythms were not significantly different between emmetropes and myopes, except for body temperature and MOPP.

Conclusions. Diurnal variations in ocular and systemic parameters were observed in young adults; however, these variations were not associated with habitual light exposure. Emmetropic and myopic refractive error groups showed small but significant differences in body temperature and MOPP, while other ocular and systemic patterns were similar.

Keywords: circadian rhythm, choroid, intraocular pressure, myopia, melatonin

Diurnal rhythms are variations in physiological parameters that are synchronized to the solar day. Diurnal rhythms in the eye have been observed in chickens,1 rodents,2 nonhuman primates,3 and humans4,5 and include variations in axial length,4 intraocular pressure (IOP),6 and choroidal thickness.4,7 In both animals and humans, eyes exposed to regular light/dark patterns undergo a decrease in axial length and increase in choroidal thickness at night, with the highest daily IOP exhibited in the morning.4,8,9 Several other physiological parameters in the body also show diurnal variations, including systemic melatonin secretion,10–12 core body temperature,13 and heart rate.14 These rhythms are mediated by light exposure and endogenous clock signals, which influence the amplitude and phase in the underlying circadian oscillator.15,16

The intrinsically photosensitive retinal ganglion cells (ipRGCs) are inner retinal cells involved in light-mediated circadian entrainment.17,18 The ipRGCs are known to influence systemic melatonin concentration in a pathway that leads from the retinohypothalamic tract to the suprachiasmatic nucleus, and, ultimately, to the pineal gland.17,19,20 A previous study1 showed that light exposure is associated with the ipRGC-driven pupil response. Additionally, known relationships exist between light exposure and diurnal rhythms of melatonin synthesis,22 and these rhythms might differ between myopic and nonmyopic individuals. Kearney et al.23 have shown that young adult myopes have significantly higher morning serum melatonin concentration than nonmyopes.

Findings in animal models suggest that light exposure patterns affect eye development and refraction. When chicks are reared in constant light or less than 4 hours of darkness per day, altered refractive errors are observed.24–27 Light exposure patterns might also differ between myopic and nonmyopic young adults. For example, a study that investigated young adult third-year law students reports an association between myopia and less daily exposure to darkness, leading investigators to suggest that perturbations in the daily light/dark cycle may also lead to refractive errors in humans.28

Accumulating evidence suggests that light exposure may influence refractive development in children. A meta-analysis29 has concluded that increased outdoor time is effective in preventing the onset of myopia. Recent longitudinal intervention studies in Eastern Asia have shown that children with increased time outdoors have reduced incidence rate of myopia30 and decreased axial growth.31,32 The underlying mechanism of protective effects of time outdoors has yet to be elucidated. Potential contributors include changes in optical factors that occur outdoors compared to indoors, such as a smaller pupil and flatter dioptric space,33 or changes in biochemical pathways, such as an increase in retinal dopamine with increased light exposure, as demonstrated in animal eyes.
Evidence suggests that diurnal variations in IOP differ with axial length and refraction. In young adults, diurnal IOP fluctuation has been shown to negatively correlate to axial length, with shorter eyes demonstrating larger IOP variation over a 24-hour period. Another study in adults reports a significant difference in the timing and amplitude of diurnal IOP variations between emmetropes and moderate to severe myopes. However, studies assessing the relationship of IOP and refraction have been conflicting, with some studies finding that myopes have higher daytime IOP than nonmyopes, and others finding no difference in IOP between refractive error groups.

This study addressed diurnal changes in young adults with and without myopia. Studying diurnal patterns in this age group is valuable for a number of reasons. Firstly, myopia carries with it a significant disease burden, including increased risks of retinal detachment and glaucoma. Understanding the diurnal changes in these parameters and their interactions, especially IOP and ocular perfusion, may improve understanding of the disease risks. Secondly, such study may provide understanding of myopia itself. Differences in diurnal changes in perfusion, IOP, light exposure profiles, and associated hormone levels may be present in this age group, as suggested in previous studies, either as a cause or consequence of myopia. Therefore, in light of reports that systemic melatonin concentration and IOP diurnal rhythms vary between myopic and nonmyopic adults, and evidence that outdoor time influences refractive development in children, we hypothesized that objectively measured light exposure is associated with ocular and systemic diurnal rhythms in young adults, and these rhythms may vary with refractive status. We measured diurnal changes in ocular and systemic parameters over a 24-hour period in emmetropic and myopic adults. Analysis of ocular biometric rhythms has been previously reported for these subjects.

Here, we present analysis of additional ocular parameters, including IOP and mean ocular perfusion pressure (MOPP), in relation to systemic physiological rhythms and objectively measured light exposure.

**Methods**

Healthy subjects, ages 22 to 41 years, participated (n = 42). Subjects provided informed consent after the purpose of the study and the risks were explained. The study was approved by the Committee for Protection of Human Subjects at the University of Houston and followed the tenets of the Declaration of Helsinki.

Subjects underwent a screening to determine ocular and systemic health. All subjects had best-corrected visual acuity of 20/20 or better. Exclusion criteria included ocular disease, the use of melatonin or other pharmacologic sleep aids, and travel outside of two time zones in the month before the experiment. Noncycloplegic autorefraction (WAM-5500; Grand Seiko, AIT Industries, Bensenville, IL, USA) was performed, and subjects were classified as emmetropic (spherical equivalent refraction [SER] of +1.50 to ≥–0.75) or myopic (SER ≤–0.75).

Following screening and enrollment, an Actiwatch Spectrum (Philips, Respironics, Bend, OR, USA) was dispensed for 1 week of continuous wear for objective measurements of each subject’s habitual light exposure, sleep, and physical activity. The light sensor in the Actiwatch measures the illuminance of broadband light in lux (range, 0.1–200,000 lux), and the irradiance of the blue (400–500 nm), green (500–600 nm), and red (600–700 nm) components (µW/cm²). Actiwatches were analyzed by using the device software, Philips Actiware 6.0.8. Activity and sleep data were evaluated to ensure that subjects had normal sleep/wake patterns. Time outdoors per day was calculated as minutes spent exposed to greater than 1000 lux. Average daily white light exposure and time outdoors for the week before the lab visit were calculated.

After 1 week, subjects presented to the lab at 8:00 AM for the experimental measurements, which were collected every 4 hours for 24 hours beginning at 8:00 AM, as described previously. Subjects were asked to refrain from caffeine, alcohol, and vigorous physical activity during the experiment, as these have been shown to affect choroidal thickness, IOP, heart rate, and body temperature. Measurements took 15 to 20 minutes at each time point. Subjects went about their activities during the day, and remained in the lab with all lights off from 11:00 PM to 7:00 AM, during which time they were encouraged to sleep. For the two time points during the night (12:00 AM and 4:00 AM), a dim red light was used for illumination, and the brightness of instrument monitors was decreased to minimize disruptions to circadian rhythms. A representative actigraph trace is shown for one subject during the 24-hour experimental period in which the subject slept in the lab to illustrate the protocol (Fig. 1).

At each time point except during the lights off period, subjects first rinsed their mouth with water in preparation for
saliva collection, and then underwent a distance-viewing period for 10 minutes while sitting upright in a chair to minimize the influence of previous activity, relax accommodation, and standardize the conditions under which ocular imaging was performed. During the distance-viewing period, subjects watched a television at 4 meters, binocularly and with habitual distance correction, when necessary.

All measurements were collected in a sitting position. Subjects were asked to collect 1 mL saliva into a vial, which was immediately placed in a -20°C freezer for analysis using melatonin and cortisol ELISA kits (Salimetrics, LLC, State College, PA, USA), as previously described. Samples were run in duplicate within 1 month of collection (stable at -20°C for up to 6 months per Salimetrics). Body temperature was measured three times at each time point by using an underwater digital thermometer with disposable probe cover (Welch Allyn, Skaneateles Falls, NY, USA). Blood pressure and heart rate were measured three times by using an electronic cuff (Omron, Bannockburn, IL, USA).

Ocular measurements were performed on the right eye only and included IOP, retinal imaging, and biometry. IOP was measured by using a rebound tonometer (iCare, Tiolat Oy, Helsinki, Finland), and three measurements, each an average of six readings, were recorded. From the diastolic (DBP) and systolic (SBP) blood pressure and IOP, MOPP was calculated for each time point, using Equation 1:

\[
MOPP = \frac{2}{3} [DBP + 1/3(SBP - DBP)] - IOP. \tag{1}
\]

Ocular imaging was performed with spectral-domain optical coherence tomography (SD-OCT; Spectralis, Heidelberg, Germany) using enhanced depth-imaging mode. At each time point, two high-quality images of the back of the eye were captured. The scan protocol included a six-line 30° radial scan centered at the fovea, with 16 frame B-scan averaging. The first image at the first time point (8:00 AM) was set as the reference for each subject, and the instrument's tracking function was used for subsequent imaging. Lastly, ocular biometry (LenStar; Haag-Streit, K¨oniz, Switzerland) was measured. Five measurements were recorded and averaged at each time point.

Analysis

Raw OCT data (*.vol files) were exported and analyzed with custom-written software in MatLab (MathWorks, Inc., Natick, MA, USA) using a semiautomated process. Data were adjusted for lateral magnification by constructing a three-surface schematic eye, as described by Bennett and Rabbets and Bennett et al. for each subject by using their measured axial length and corneal curvature. Individualized transverse scaling was then calculated, assuming a spherical retina as previously described. Bruch’s membrane, the inner limiting membrane, and the retinal layers, including inner and outer photoreceptors, were automatically segmented and manually corrected when necessary. Image contrast was optimized, and the sclera/choroid border was segmented manually. Axial choroidal thickness was calculated as the distance from Bruch’s membrane to the posterior choroid/sclera border for 1536 points along each of the six scan lines. Data were binned for the central 1-mm diameter and averaged for the two radial scans centered on the fovea of the right eye, which were collected at each time point. Repeatability and retinal and choroidal thicknesses in the 3-mm and 6-mm annuli by quadrant are presented in detail elsewhere.

Statistical analyses were performed by using MedCalc Software (12.5.3.0; MedCalc, Mariakerke, Belgium) and Excel (Microsoft Office 2013; Microsoft, Redmond, WA, USA). On the basis of data from our previous investigations, with a between-groups difference of 59 μm, a within-subjects standard deviation of 75 μm, and a cosine change in choroidal thickness across the day of 9.6-μm amplitude, 15 subjects per group would provide power to measure a time-of-day effect at alpha < 0.05. Data are expressed as mean ± standard error unless otherwise noted. Normality was assessed with the Shapiro-Wilk test. A critical value <0.05 was considered statistically significant. For each parameter for each subject, the mean of the values from all time points was calculated, and the amplitude of diurnal variation was determined as the difference between the maximum (peak) and minimum (trough) values across 24 hours. To investigate whether significant diurnal changes occurred over a 24-hour period, a repeated measures analysis of variance (ANOVA) with one within-subject factor (time of day) and one between-subjects factor (refractive error group) was used. The ANOVA was used to detect whether average measurements changed across the day for different times and between groups, and is a sensitive test if amplitudes and phase do not show large within-group variation. However, ANOVAs do not represent the most appropriate test for assessing amplitude and phase independently. For example, groups might have different amplitudes of change across the day; however, if there is a large variation in phase within groups (i.e., variation in peak phase between subjects) then the ANOVA would not yield significant results. Likewise, there might be differences in acrophase (the time of maximum response) between groups, but because of variation between participants’ amplitude, the ANOVA may not detect between-group differences in acrophase. Therefore, additional analysis was undertaken to assess whether amplitude (ignoring phase) differed between groups, and to see whether acrophase (ignoring amplitude) differed between groups. To estimate the amplitude and acrophase of diurnal rhythm for each parameter for each subject, Fourier analysis was used to determine the fundamental cosine, using Equation 2.

\[
y = \text{Diurnal Mean} + \frac{\text{Amplitude}}{2} \cos\left(\frac{2\pi}{24}(t - \text{Acrophase})\right). \tag{2}
\]

where \(t\) is time of measurement (on a 24-hour clock), \(\text{Acrophase}\) is the time where the fitted cosine reaches its peak, and \(\text{Amplitude}\) is the difference between maximum and minimum \(y\) values in the fitted cosine.

Mann-Whitney \(U\) tests were used to analyze refractive error group differences in the amplitude of diurnal variation. Group average peak acrophase was estimated as the vector mean of acrophases distributed around the unit circle. Rayleigh tests were used to investigate whether the acrophase distribution was significantly different from a uniform distribution across the day. As part of the Rayleigh test calculation, \(r^2\), an estimate of the dispersion of acrophase across the day, was estimated. An \(r^2 = 0\) indicates that acrophase is distributed evenly across the day. Higher values of \(r^2\) indicate clustering of acrophase at a given time, with \(r^2 = 1\) occurring when all acrophases occur at the same time. Watson’s \(U^2\) test was used to assess whether there was a significant diurnal difference between myopes and emmetropes in terms of acrophases. For parameters that showed a refractive error difference, data were fit with cosine functions to compare rhythms. Pearson correlation was used to investigate relationships between the amplitude of diurnal variation for each parameter and mean daily light exposure.

Results

Subject demographics and ocular characteristics are shown in Table 1. Mean subject age was 27.2 ± 4.2 years, with 14 males and 28 females. The SER of all right eyes was -2.55 ± 3.13
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#### DISCUSSION

Ocular factors, including the choroid, IOP, and perfusion pressure, and systemic factors, including sleep and stress hormones and body temperature, have potential roles in refractive development, and valuable information can be gained by investigating how these parameters fluctuate across the day in refractive groups. This study examined diurnal rhythms of ocular and systemic processes, and extended...
### Table 2: Objectively Measured Mean (± Standard Error) Daily Time Outdoors, Daily Light Exposure, Nightly Light Exposure, Sleep Duration, and Physical Activity for the Emmetropic and Myopic Groups

| Parameter                              | Total          | Emmetropes  | Myopes       | P Value |
|----------------------------------------|----------------|-------------|--------------|---------|
| Time outdoors, min                     | 92.2 ± 7.33    | 103.2 ± 12.91 | 82.59 ± 8.29 | 0.17    |
| Daily white light exposure, lux        | 1130.82 ± 110.15 | 1152.28 ± 164.74 | 1115.61 ± 150.51 | 0.87    |
| Nightly white light exposure, lux      | 1.31 ± 0.2     | 1.54 ± 0.41  | 1.28 ± 0.2   | 0.89    |
| Sleep duration, min                    | 442.1 ± 7.9    | 438.44 ± 13.31 | 444.65 ± 9.87 | 0.70    |
| Sleep efficiency, %                    | 89.5 ± 0.54    | 90.03 ± 0.9  | 89.13 ± 0.2  | 0.42    |
| Physical activity, counts/min         | 264.81 ± 9.4   | 268.67 ± 16.46 | 262.08 ± 11.33 | 0.75    |

P value for unpaired t-test.

### Table 3: Daily Mean (± Standard Error) for Each Parameter for All Subjects, for Emmetropes, and for Myopes

| Parameter                              | All            | Emmetropes | Myopes       | P Value |
|----------------------------------------|----------------|------------|--------------|---------|
| Axial length, mm                       | 24.66 ± 0.23   | 23.77 ± 0.25 | 25.26 ± 0.30 | <0.001* |
| Choroidal thickness, central 1 mm, µm  | 334.62 ± 12.28 | 368.33 ± 17.72 | 305.93 ± 14.5 | 0.009*  |
| Intraocular pressure, mm Hg           | 14.15 ± 0.42   | 14.20 ± 0.61  | 14.12 ± 0.58  | 0.71    |
| Body temperature, °C                   | 36.87 ± 0.04   | 36.86 ± 0.07  | 36.88 ± 0.05  | 0.95    |
| Heart rate, bpm                        | 72.63 ± 1.58   | 71.52 ± 2.54  | 73.39 ± 2.05  | 0.80    |
| Mean arterial pressure, mm Hg         | 83.43 ± 0.88   | 83.46 ± 1.29  | 83.42 ± 1.20  | 0.95    |
| Mean ocular perfusion pressure, mm Hg | 41.47 ± 0.73   | 41.44 ± 0.95  | 41.49 ± 0.98  | 0.95    |
| Melatonin, pg/mL                       | 11.72 ± 0.78   | 10.64 ± 1.22  | 12.41 ± 1.0   | 0.25    |
| Cortisol, dg/mL                        | 0.23 ± 0.01    | 0.25 ± 0.02   | 0.21 ± 0.01   | 0.18    |

P values from Mann-Whitney U test.
* Significance at P < 0.05.

### Table 4: Amplitude of Diurnal Variation (Mean ± Standard Error) and Acrophase for Each Measured Parameter for All Subjects (n = 42)

| Parameter                              | Amplitude | Acrophase, Peak, h | Time of Day | Refractive Error | Time by Refractive Error |
|----------------------------------------|-----------|--------------------|-------------|------------------|--------------------------|
| Axial length, µm                       | 35.71 ± 2.99 | 12.96              | <0.001*     | 1.00             | 0.501                    |
| Choroidal thickness, central 1 mm, µm  | 25.80 ± 2.08 | 11.42              | <0.001*     | 0.96             | 0.42                     |
| Intraocular pressure, mm Hg           | 4.92 ± 0.25  | 15.9               | <0.001*     | 0.96             | 0.42                     |
| Body temperature, °C                   | 0.52 ± 0.04  | 21.66              | 0.034*      | 0.11             | 0.75                     |
| Heart rate, bpm                        | 16.38 ± 0.88 | 17.97              | <0.001*     | 0.14             | 0.71                     |
| Systolic blood pressure, mm Hg        | 11.45 ± 0.7  | 20.85              | 0.015*      | 0.94             | 0.70                     |
| Diastolic blood pressure, mm Hg       | 11.89 ± 0.67 | 19.14              | <0.001*     | 0.36             | 0.88                     |
| Mean arterial pressure, mm Hg         | 9.70 ± 0.49  | 22.02              | <0.001*     | 0.88             | 0.55                     |
| Mean ocular perfusion pressure, mm Hg | 26.95 ± 1.88 | 3.19               | <0.001*     | 0.69             | 0.19                     |

P values are from two-factor repeated measures ANOVA for time of day, refractive error, and time of day by refractive error.
* Significance at P < 0.05.

![Figure 2](https://example.com/figure2.png)

**Figure 2.** Mean (± standard error) 24-hour change in body temperature (°C) for emmetropes (filled symbols) and myopes (open symbols), lines are cosine fits to the data, gray area represents the dark period.
previous work to investigate potential association with light exposure and refractive status in young adults. Our results showed that there were significant diurnal variations in ocular parameters, including axial length, choroidal thickness, IOP, and MOPP, as well as systemic parameters, including blood pressure and heart rate, body temperature, and melatonin and cortisol levels, consistent with previous reports. Furthermore, we showed small but significant differences in diurnal rhythms of body temperature and MOPP between emmetropic and myopic groups.

**FIGURE 3.** Diurnal amplitudes and acrophase (hours) calculated by Fourier analysis (Equation 2). (A) Axial length (µm), (B) choroidal thickness (µm), (C) IOP (mm Hg), (D) MOPP (mm Hg), (E) body temperature (°C), (F) mean arterial pressure (mm Hg), (G) melatonin concentration (pg/mL), (H) cortisol concentration (dg/mL). Median acrophases are indicated, calculated by averaging unit vectors. Emmetropes, filled circles; myopes, unfilled circles.
For all parameters investigated, the amplitudes of diurnal variations were not associated with light exposure during the previous week. We made use of an ambulatory, noninvasive device, the Actiwatch, for measures of habitual light exposure and sleep patterns before the experimental visit. We found that time per day exposed to outdoor light (>1000 lux) in our young adult emmetropic group was not significantly different from that of the myopic group (103 and 82 minutes, respectively). This is similar to findings in previous reports using objective methods to measure light exposure in young adult populations, which range from 74 to 112 minutes per day of exposure to outdoor light levels. These latter young adult populations were also drawn from university settings.

In accordance with previous reports, diurnal variations in IOP and axial length were in phase with each other and demonstrated a weak but significant correlation ($R^2 = 0.11$); as IOP increased, axial length increased. It has been proposed that increased IOP may induce axial length increases in myopia. Evidence from studies in chicks suggests that an increase in IOP can lead to biomechanical changes in the sclera that result in an increase in axial length. Similarly, studies in humans who have undergone glaucoma filtration surgery to lower IOP have found decreases in axial length. However, increasing IOP does not result in an increase in axial length of the tree shrew, whose sclera is structurally more similar to that of humans than of the chick. Additionally, it is possible to dissociate IOP and axial length rhythms in chicks. These latter findings suggest that increases in axial length are not a result of biomechanical changes due to increases in IOP.

Previous studies assessing diurnal variations in IOP with refractive error show conflicting results, with some demonstrating that patterns in IOP rhythms are different between emmetropic and myopic subjects, and others showing no difference. Liu et al. have collected 24-hour data of IOP, axial length, blood pressure, and heart rate in subjects with either emmetropic or moderate to severe myopia. The authors have found significant differences in the magnitude of nighttime increases in IOP with refractive error, as well as timing for supine IOP measurements. The mechanism behind the refractive error group differences is unclear, although the authors speculate that choroidal vascular volume and episcleral pressure may play a role. Additionally, studies have shown that myopic scleras are biomechanically weaker than emmetropic scleras, and more susceptible to axial elongation with increased short-term IOP elevation. Here, we found that diurnal patterns in IOP were not significantly different between emmetropes and myopes, similar to a report by Chakraborty et al. While qualitative differences can be observed in the cosine fits for each group (Fig. 5), with the myopic group tending to have a smaller amplitude in diurnal variation, similar to results reported by Liu et al., differences did not reach statistical significance. Both of the previous studies were carried out in a young adult population of emmetropes and myopes, similar to the current study. Differing results between studies could be due to differences in how IOP was measured; our measurements were recorded in a seated position, while Liu et al. have found differences between groups when IOP is recorded in a supine position.

Myopia has been shown to carry an increased risk of primary open angle glaucoma. IOP and MOPP are well known to be associated with glaucoma. Ocular perfusion pressure is the pressure available to drive blood through the intraocular vasculature. Larger 24-hour fluctuations in MOPP have been shown to be a risk factor for glaucoma severity in normal tension glaucoma. Here, Rayleigh test for acrophase detected a small but significant difference in the acrophase of MOPP between emmetropes and myopes, while amplitude was not significantly different between groups. It is uncertain if this observed phase shift is a clinically meaningful result, and whether it has implications in ocular perfusion.

With known relationships between light exposure and melatonin release, and increasing evidence that light exposure has a protective effect for myopia, speculation exists whether melatonin may play a role in refractive error development. Melatonin is a hormone that is released in darkness from the pineal gland.
In the retina, melatonin has a reciprocal relationship with dopamine, a neurotransmitter linked to regulate eye growth, as demonstrated in animal models. Systemic melatonin production is regulated by the suprachiasmatic nucleus and plays a significant role in regulating sleep/wake patterns. Recent studies have suggested that sleep patterns may vary between nonmyopic and myopic individuals, with myopes reporting poorer sleep quality compared to nonmyopes. Ayaki et al. have found decreased sleep quality in myopic children, with no differences in sleep between adult myopes and nonmyopes. Here, we found that diurnal variations in salivary melatonin concentration were similar between emmetropic and myopic subjects. Our subject population was from a group of young adults, drawn from students, faculty, and staff at the university. Subjects demonstrated similar sleep/wake cycles and light exposure patterns, as measured objectively with a wrist-worn sensor during the week previous to diurnal measurements. We speculate that evaluating subjects with a wider range of light exposure patterns may reveal associations between light exposure and diurnal rhythms.

Cortisol is a hormone that is associated with waking, alertness, and stress, and is known to exhibit a diurnal rhythm that peaks in the early morning. While melatonin synthesis is known to be light dependent, cortisol has been shown to be regulated internally and remain phase locked to melatonin rhythms. Therefore, cortisol is considered to be a reliable marker of circadian rhythm. Similar to findings for melatonin concentration, the diurnal pattern in cortisol concentration observed here was not significantly different between refractive error groups, with a peak cortisol concentration at 8.86 hours, shortly after the dark period ended.

Body temperature is a well-described, simple, and noninvasive marker of a systemic diurnal rhythm. Physical activity is known to elevate body temperature and modify an individual’s capacity for thermoregulation, an effect that can persist for hours after activity has ceased. For these reasons, it was of interest to assess body temperature in this study. We found that diurnal variations in body temperature were significantly different for myopic and emmetropic subjects (two-factor repeated measures ANOVA), with myopic subjects exhibiting less diurnal variation than emmetropes. Body temperature is known to undergo diurnal variations with the lowest temperature in the early morning and the highest temperature in the late evening. Many endogenous and exogenous factors contribute to body temperature, including hormones, fitness level, age, diet, and lifestyle. Emmetropic and myopic groups in this study did not show a difference in objectively measured physical activity levels. It is unclear why a difference was observed between refractive error groups. Future investigations may be able to provide insight into this unexpected finding.

Studies have shown that changes in ocular growth patterns are induced in animal models, disruptions in diurnal rhythms are observed. For example, in chick eyes undergoing decreased growth through induced myopic defocus, the rhythm in axial length phase-delays, while the rhythm in choroidal thickness phase-advances, bringing the two into phase with each other. It is likely that we failed to detect a difference between refractive error groups because the subjects in our myopic group were stable myopes, whereas in chick studies, the animals were undergoing active eye growth and myopia development when altered rhythms were observed. To address this difference, studies similar to this one, carried out in children or young adults who are undergoing myopia development and progression, will be important in clarifying the relationship of axial length and choroidal thickness rhythms in eye growth.

The study design used here presented some limitations. While all subjects demonstrated regular sleep/wake patterns and similar light exposure as recorded by the Actiwatch, it is possible that behaviors were influenced by virtue of being observed; a longer observation period or administration of behavioral questionnaires may have revealed variations in behaviors that might influence diurnal rhythms. During the experimental measures in lab, it was necessary to wake subjects twice during the night and use dim illumination for measurements. A previous study has shown that brief periods of moderate illumination during the night do not interrupt diurnal variations in IOP; it is unlikely that the dim red illumination used here altered diurnal rhythms. Additionally, all measurements were recorded with the subjects in a seated position, as opposed to a more natural prone or supine position during the night. However, a study has shown that IOP diurnal variations are observable for both sitting and supine positions, with no significant differences in the

**Figure 6.** Mean (± standard error) change in melatonin concentration (emmetropes, filled circles; myopes, open circles) and cortisol concentration (emmetropes, filled squares; myopes, open squares) across 24 hours; gray area represents the dark period.
amplitude based on position. Another study has shown that choroid volume does not differ in subjects between recumbent and sitting positions. It is unlikely that a change in position would have altered observed diurnal rhythms. Finally, subjects’ refraction was only measured on one occasion, so we were unable to assess whether myopic subjects were progressing or stable myopes. However, subjects were likely to be stable myopes, given their age range of 22 to 41 years, which is outside the range over which juvenile-onset myopia typically progresses.

In summary, we described diurnal variations in ocular and systemic parameters over a 24-hour period. Emmetropic and myopic refractive error groups showed small but significant differences in body temperature and MOPP rhythms, while other ocular and systemic patterns were similar. Previous light exposure was not associated with the amplitude of diurnal variation in axial length, choroidal thickness, MOPP, body temperature, melatonin concentration, or cortisol concentration in this group of young adults.

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