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The post illumination pupil response is reduced in seasonal affective disorder

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Abstract

Individuals with seasonal affective disorder (SAD) may have a decreased retinal sensitivity in the non-image forming light-input pathway. We examined the post illumination pupil response (PIPR) among individuals with SAD and healthy controls to identify possible differences in the melanopsin signaling pathway. We also investigated whether melanopsin gene (*OPN4*) variations would predict variability in the PIPR. Fifteen SAD and 15 control participants (80% women, mean age 36.7 years, *SD* = 14.5) were assessed in the fall/winter. Participants were diagnosed based on DSM-IV-TR criteria. Infrared pupillometry was used to measure pupil diameter prior to, during, and after red and blue stimuli. In response to blue light, the SAD group had a reduced PIPR and a lower PIPR percent change relative to controls. The PIPR after the blue stimulus also varied on the basis of *OPN4* I394T genotype, but not *OPN4* P10L genotype. These findings may indicate that individuals with SAD have a less sensitive light input pathway as measured by the PIPR, leading to differences in neurobiological and behavioral responses such as alertness, circadian photoentrainment, and melatonin release. In addition, this sensitivity may vary based on sequence variations in *OPN4*, although a larger sample and replication is needed.

Keywords

Seasonal affective disorder; melanopsin; pupillometry

1. Introduction

Seasonal affective disorder (SAD) accounts for approximately 10–20% of outpatients with recurrent depression (Magnusson and Partonen, 2005). Treatment of this large segment of depressed individuals may be improved with a better understanding of the etiology of SAD. Although the pathophysiologic mechanisms underlying SAD remain unknown, one hypothesis postulates a seasonal variation in retinal subsensitivity (Reme et al., 1990; Hebert

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et al., 2002). Retinal subsensitivity to light is suggested by many of the phenomena observed in SAD including: the regular winter timing of depressive episodes, a lengthened melatonin release profile during longer winter nights (Wehr et al., 2001), the utility of light therapy as an antidepressant treatment (Rohan, et al., 2009), and retinal electrophysiology research (reviewed below). Individual differences in retinal signaling may explain retinal subsensitivity (Wehr et al., 2001). Specifically, retinal subsensitivity in SAD may result from the absence of a seasonal change in retinal sensitivity found in healthy controls, such that the retina fails to become more sensitive to the low light levels of winter. In contrast to previous studies, the pupillometry method in the present study is designed to isolate the response of melanopsin-containing, intrinsically photosensitive retinal ganglion cells (ipRGC), which are the primary class of photoreceptors entraining the circadian clock. A polymorphism in the melanopsin gene has been associated with increased risk of SAD, suggesting that the melanopsin gene may be etiologically significant in SAD (Roeklein et al., 2009).

In addition, melanopsin-driven processes have significant overlap with SAD symptomatology. Melanopsin cells in the retina project to multiple areas in mammals (Do and Yau, 2010), and if these projections are conserved in humans, they would include areas involved in circadian regulation (suprachiasmatic nuclei, intergeniculate leaflet), energy homeostasis (lateral hypothalamus), and sleep regulation (ventral subparaventricular zone, preoptic nucleus; Hattar et al., 2002; Hannibal and Fahrenkrug, 2004; Hattar et al., 2006). Circadian, energy, and sleep processes are either involved in the symptoms of SAD (e.g., fatigue), or hypothesized to be etiologically significant in SAD.

Given the overlap between SAD symptomatology and melanopsin-driven processes, we hypothesized that retinal subsensitivity in SAD may be mediated at least in part by melanopsin containing ipRGCs. Theoretically the ipRGC sensitivity in SAD could be lower than that of healthy individuals, such that the environmental irradiance levels in winter may fall below this threshold, leading to depression. The present study measures melanopsin cell sensitivity in individuals with SAD compared to non-depressed controls.

Melanopsin containing ipRGCs project from the retina to the suprachiasmatic nucleus, which serves as a circadian clock, and have a peak sensitivity of ~482 nm when measured in vitro (Provencio et al., 2001; Lucas et al., 2001; Hattar et al., 2002; Berson et al., 2002; Gooley et al., 2003; Dacey et al., 2005) and peak sensitivity of 482 nm when measured in vivo (Gamlin et al., 2007; Markwell et al., 2010). In addition to entraining the circadian clock, ipRGCs are involved in the pupil light reflex, in which the pupil constricts and redilates in response to light. The pupil area in melanopsin knockout mice after 1 min of blue light is much larger (i.e., less constricted) than that of wild-type mice (Hattar et al., 2003; Lucas et al., 2003). Using primate cell recording techniques, Gamlin et al. (2007) and Dacey et al. (2005) demonstrated that, unlike rods and cones, ipRGCs have a sustained response after removal of the light source. This results in continuing constriction of the pupil after light off and is called the post-illumination pupil response (PIPR). The PIPR is intrinsic to melanopsin-containing ipRGCs (Gamlin et al., 2007). Therefore, the magnitude of the PIPR response to blue light reflects the sensitivity of the melanopsin pathway (Gamlin et al., 2007). The PIPR then provides an elegant way to isolate the contributions of ipRGCs from those of rods and cones. The PIPR has been measured in a range of healthy and clinical populations (Gamlin et al., 2007; Kawasaki et al., 2007; Kardon et al., 2009; Kankipati et al., 2010; McDougal et al., 2010; Kardon et al., 2011; Kankipati et al., 2011; Feigl et al., 2011; Park et al., 2011; Kawasaki et al., 2012; Feigl et al., 2012), and the present design is based in part on the methods in these studies. However, the PIPR has yet to be measured in SAD.

Electroretinography (ERG) is another tool that has been used to assess retinal sensitivity in SAD. ERG records electrical responses at the corneal surface to measure the function of retinal cells such as rods (scotopic) or cones (photopic), but not retinal ganglion cells. Photoreceptor (and bipolar cell) sensitivity is assessed by the measure log K, which is a calculation of the intensity of light that evokes half of the maximum (V_{max}) rod or cone response. Using this measure, Lavoie et al. (2009) found that rods were less sensitive in SAD patients compared to controls during winter measurements. They also showed that the V_{max} amplitudes of both rods and cones were reduced in SAD patients compared to controls. Most interestingly, all three of these measures were normalized in SAD following light therapy treatment (Lavoie et al., 2009). Decreased rod sensitivity seems to be a state marker for depression in SAD, as Gagne and Hebert (2011) also found that rods had decreased sensitivity in SAD during winter, which normalized during the summer. While studying the effects of light history on retinal sensitivity, Gagne and Hebert (2011) identified a possible trait marker for SAD. In participants with SAD, exposure to bright light for 1 hour prior to scotopic measurement caused a decrease in V_{max} during winter and summer measurements (Gagne and Hebert, 2011). Because ipRGCs are most sensitive to blue light, Gagne et al., (2011) compared retinal responses of SAD and control participants after exposure to red or blue light. In response to blue light, both groups had a decrease in an ERG measure that is driven by ON-bipolar cells and Muller cells. These data reviewed here support a role for retinal subsensitivity in SAD, but do not assess the specific contribution of ipRGCs. Given the relationship between SAD, retinal subsensitivity, and gene polymorphisms in melanopsin, we hypothesize that ipRGCs specifically will be less sensitive in SAD participants than in controls when measured using a unique pupillometry measure specific to ipRGCs.

The present study was designed to specifically test the sensitivity of ipRGCs in SAD using a melanopsin-specific pupillometry test. M1-type melanopsin cells are directly involved in the pupillary light reflex as well as in circadian photoentrainment (Schmidt et al., 2011). Hence, impairment in one process (i.e., pupil constriction) may reflect impairment in the other processes mediated by M1 cells (i.e., photoentrainment). Although rods and cones also contribute to circadian photoentrainment (Provencio, 2011), melanopsin cells drive 40–50% of the response (Do and Yau, 2010). Because melanopsin cells are such a large component of the pathway implicated in retinal subsensitivity in SAD, it is important to investigate whether there may be a decrease in melanopsin cell sensitivity. Therefore, in the present study, we compared individuals with SAD and controls on the melanopsin-driven PIPR. We also examined differences in the PIPR by melanopsin gene (*OPN4*) sequence variations. Although others (i.e., Higuchi et al., 2013) have found that the steady-state pupil diameter varies as a function of melanopsin gene variation I394T, this will be the first test of the melanopsin-specific PIPR in SAD with both the P10L and I394T melanopsin gene variations.

2. Methods

2.1 Participants

Participants age 18–65 were recruited from the greater Pittsburgh, Pennsylvania metropolitan area through community advertising (latitude 40°26 N). The institutional review board of the University of Pittsburgh approved this study, and participants gave informed consent and received compensation. None of the participants reported a history of psychotic or bipolar disorder, sleep disordered breathing, narcolepsy, or current substance use disorder. All participants were studied during the fall or winter months in 2011 and 2012 while individuals in the SAD group were in a Major Depressive Episode.

2.2 Clinical Assessments

SAD group participants met Diagnostic and Statistical Manual of Mental Disorders (DSM-IV-TR; APA, 2000) criteria for unipolar Major Depressive Disorder With a Seasonal Pattern according to the Structured Clinical Interview for DSM-IV Axis I Disorders - Research Version (SCID-I; First et al., 2002). Participants also completed the Modified Seasonal Pattern Assessment Questionnaire (M-SPAQ; Blouin et al., 1992; Rosenthal et al., 1984), the Structured Interview Guide for the Hamilton Depression Rating Scale, Seasonal Affective Disorder version (SIGH-SAD; Williams et al., 1992), and the Beck Depression Inventory, Second Edition (BDI-II; Beck et al., 1996). Inclusion criteria for the SAD group were (1) a diagnosis of MDD-SP based on the SCID, (2) a SIGH-SAD score of > 19 total, > 9 on the HAM-D subscale, and > 4 on the atypical subscale which defines an SAD episode (Terman et al., 1990), and (3) a Global Seasonality Score (GSS) from the M-SPAQ of > 11 , endorsing “feeling worst” in January and/or February (with or without other months) but not July and/or August, and perceiving seasonal changes as at least a “moderate” problem (Kasper et al., 1989). Inclusion criteria for the control group were (1) no history of mood disorder based on the SCID, (2) scores below episode criteria listed above on the SIGH-SAD, (3) a GSS of 8 or 9 and endorsing “no problems” with seasonal changes on the M-SPAQ, or GSS < 8 (Kasper et al., 1989), and (4) BDI-II score < 10 .

2.3 Stimuli

Lights of equal corneal irradiance ($13.7 \log \text{Photons/cm}^2/\text{s}$) were presented for 30s ON, and 120s OFF. Retinal irradiance was calculated to account for age-related decreases in lens transmission and pupil diameter during the stimulus. Using age and pupil diameter during stimulus presentation, retinal irradiance was calculated for each participant. Stimuli were turned on and off electronically using a commercially available remote receiver and E-Prime 2 software (Psychology Software Tools, Inc., Pittsburgh, PA). Round light emitting diode bulbs (Super Bright LEDs; St. Louis, MO) were placed behind an 8.5 diameter mylar diffuser (Speedotron, Chicago, IL) at a distance of 17 from the eye ($\sim 29^\circ$ of the visual field; lenses were not used to focus the stimulus on a point in the retina). The LED lights were presented to both eyes, and pupil responses were measured in the left eye in all subjects, so the consensual response was not used. Stimuli are 22.68 nm (467.7 nm blue light) and 15.78 nm (632.9nm red light) full-width half-maximum (FWHM), and absolute corneal irradiance was measured at the approximate location of the participants' eye in the headrest with a 50-micron width aperture spectrophotometer (USB4000 Fiber Optic Spectrometer, Ocean Optics, Dunedin, FL). Due to findings that ipRGCs may be bistable and responses to blue light could be potentiated by red light (Melyan et al., 2005; Mure et al., 2009; Mure et al., 2007), we did not randomize the order of stimuli presentation. Instead, the blue stimulus was presented first, followed by red. This pattern was repeated twice.

Calculated retinal irradiance (E_r) was averaged across all subjects for the blue and red stimuli (Blue $E_r=12.3 \log \text{Photons/cm}^2/\text{s}$; Red $E_r=12.5 \log \text{Photons/cm}^2/\text{s}$; Pokorny et al., 1987). We calculated the retinal irradiance (E_r) based on the measured corneal irradiance (E_c), the lens transmittance for the given wavelength ($T(\lambda, \text{age})=10^{-D}$; where $D(\lambda, \text{age})$ is the optical density from Pokorny 1987), the solid angular size of the light screen ($\omega_{\text{screen}}=0.378$ steradians; from the diameter of the screen, $d=6.30$ cm, which was at a distance $ds=8.66$ cm from the subject's pupil), and the solid angular size of the image on the retina (ω_{eye} ; which was calculated based on the size of the pupil aperture diameter, $a=3\text{--}5$ mm, and the typical focal length of the adult human eye, $f=17.0$ mm,).

1. $\omega_{\text{screen}} = 2 \pi (1 - \cos(\arctan(d/2) / ds)) = 0.378$ steradians
2. $\omega_{\text{eye}} = 2 \pi (1 - \cos(\arctan((a/2) / f)) = 0.03 \text{ -- } 0.05$ steradians
3. $E_r = (\omega_{\text{eye}} / \omega_{\text{screen}}) T(\lambda, \text{age}) E_c$

The minimum pupil diameter during stimulus presentation was used for the individual pupil aperture, a , for each participant, and was $M = 0.35$ mm for the blue stimulus and $M = 0.34$ mm for the red stimulus. However, individual diameter and age were used to calculate retinal irradiance for each participant.

2.4 Absence of blue light hazard

Blue wavelength light can lead to photochemical injury to the retina between 400nm – 550nm (Slone, 1994). To ensure the safety of the light stimuli, the time-integrated (60s) hazard-weighted irradiance was calculated and found to be lower than the limit established by the American Conference of Governmental Industrial Hygienists (ACGIH, 2011) for a large angle light source by a factor of 36,000.

2.5 Pupillometry

Pupil area was measured with near-infrared illumination and solid-state video pupillometry with the EyeTrac 6 at 60 Hz using an EYE-TRAC (R) 6000 processing unit (Applied Science Laboratories, Inc.; Bedford, MA). The focal length was not changed from factory setting, which was 27mm with an F-Number of 4.5 for the EyeStart camera. Testing was constrained to between 2:00 pm and 5:30 pm, as data suggest that the magnitude of the blue PIPR (Zeile et al., 2011) and iris constriction varies across the day (Figueiro et al., 2005). An 11 min dark adaptation period was used prior to the first blue stimulus, although level of dark or light adaptation is unlikely to diminish the target response because melanopsin cells show resistance to bleaching and little adaptation under continuous light (Zhu et al., 2007). Participants were tested alone in a dark room, but could communicate with experimenters during the study by intercom. Due to significant autonomic modulation of the pupil and resultant potential effects of noise or surprise on pupil constriction, experimenters refrained from contacting participants by intercom during testing. Participants were instructed to maintain fixation at a point projected on a computer screen that was visible when the stimuli were either on or off. Participants rested in a table-mounted adjustable forehead and chinrest to minimize head movement. Data were processed using EyeNal Analysis Software for the ASL EyeTrac 6 pupillometry system (Applied Science Laboratories, Inc.; Bedford, MA), and blinks were removed using custom scripts in MATLAB software (MathWorks, Inc.; Natick, MA). Prior to removing blinks, the percentage of data lost to blinks and fixation loss was calculated. Calibration of the pupillometer with known size standard stimuli yielded a scaling factor of 11.9 and an intercept of 0.181. These values were used to convert the arbitrary units of pupil diameter recorded by the ASL system to millimeters.

2.6 Genotyping

In past studies, two coding variations in *OPN4* have been tested for increased risk of SAD (Roecklein et al., 2009). These two variations are the only coding variations in *OPN4*, and are referred to as P10L (rs2675703) and I394T (rs1079610). Both single nucleotide polymorphisms (SNPs) were genotyped from saliva samples for the present study using high-resolution melt analysis based on the melting behavior of fluorescently labeled DNA (Reed et al., 2007). Due to variations in the frequency of *OPN4* alleles across ethnicities, only those of Caucasian ancestry were assessed in the present study.

2.7 Visual Information

Although not confirmed by eye or medical examination, all participants reported no history of any retinopathy, glaucoma, cataracts, amblyopia, macular degeneration, congenital color vision deficiencies, or any type of blindness (e.g., scotoma, night blindness, etc.). Individuals with corrected vision were not excluded from the study, but were encouraged to

wear contact lenses during testing if possible to minimize interference during recording from glasses frames.

2.8 Statistical Analysis

Prior to data analysis, participants with more than 10% data loss were excluded from analyses ($n = 2$). Data analysis included (1) blink removal, (2) averaging pupil diameter (mm) across epochs of testing, and (3) diagnostic and genotype group comparisons. Groups were compared on the frequency of fixation losses and blinks. We used linear interpolation to define and replace blink data points previously described by Steinhauer et al. (2000; MATLAB, Natick, MA). The baseline was defined as the average pupil diameter during the 7 seconds prior to each stimulus. Constriction during lights on was measured as the mean pupil diameter during the 30-second stimulus (i.e., mean diameter from stimulus onset to 30 seconds after onset). Sustained response was defined as the mean of the diameter starting 10 seconds after stimulus offset for 30 seconds (i.e., 10 seconds after stimulus offset to 40 seconds after stimulus offset). The re-adaptation period in dark after stimulus presentation was 2 minutes, although shorter durations have been used (Herbst et al., 2011). We calculated the following values consistent with Kankipati et al. (2010, 2011):

1. PIPR (mm) = [baseline pupil diameter mm – Sustained pupil diameter mm]
2. PIPR Change (%) = [(PIPR X 100)/Baseline]
3. Net PIPR (mm) = (Blue PIPR – Red PIPR)
4. Net PIPR Change (%) = [Blue PIPR Change (%) – Red PIPR Change (%)]

Analysis of covariance (ANCOVA) was used to compare groups on these outcome variables, including the diameter during lights on. The Composite Scale of Morningness (CSM; Smith, Reilly, & Midkiff, 1989) was used to estimate chronotype for each person. Variables that may be associated with the PIPR or diagnosis such as chronotype (CSM score), time of awakening, calculated retinal irradiance, time of day at assessment, age, gender, and calculated retinal irradiance were tested for correlation with outcome variables, group association, and were included as covariates in ANCOVA if significantly associated (only age and gender were ultimately retained as covariates). All ANCOVAs were conducted using SPSS Statistics for Macintosh, version 19.0 (SPSS Inc., Chicago, IL). Summary data are reported as mean (M) \pm standard deviation (SD). All tests were two-tailed, and a P value of 0.05 was used.

Our first study hypothesis was that groups would not differ on variables expected to influence retinal irradiance including age, baseline pupil diameter, pupil diameter during the stimulus presentation, and calculated retinal irradiance. Second, we tested to be sure groups did not differ on factors that could affect the sensitivity of the melanopsin system such as time of assessment, daylength on the day of testing, and individual chronotype. Assuming that those factors were equally distributed across the SAD and control groups, our third and main hypothesis was that individuals in the SAD group would demonstrate a less robust PIPR in response to the blue test stimulus compared to the control group, but that no differences between groups in response to the red control stimulus would be observed.

3. Results

Groups did not differ based on age ($F_{1,29} = 0.058$, $P = 0.81$), or gender ($X^2_{1,29} = 0.995$, $P = 0.32$). Participants ranged from age 20 to 60, and the mean and standard deviation of age by group are reported in Table 1. In addition, age, gender, daylength on the testing day, time of testing, and calculated retinal irradiance were not significantly correlated with the main outcome variable, the blue PIPR (all P 's > 0.05 ; see Table 1) or associated with group

membership (see Table 1). Age and gender were retained as covariates in the main analyses. The percentage of data lost due to fixation losses and blinks was calculated, and did not differ between groups (SAD: $M = 12\%$, $SD = 8\%$; Control: $M = 10\%$, $SD = 7\%$; $F(1, 29) = .42$, $p = .53$).

Although chronotype as measured by the CSM was not associated with the PIPR or with group membership (p 's $> .05$), individuals in the SAD group had lower scores indicating a trend towards evening chronotype ($M = 32.3$) and controls had a higher mean indicating a relative trend towards morning chronotype ($M = 36.9$). However, both groups fall in the "intermediate chronotype" scoring range (i.e., 27–41) on the CSM. Individuals did differ based on self-reported time of awakening ($F_{1,29} = 4.29$, $P = 0.048$), with the SAD group getting up later on average ($M = 8:09$ AM) compared to the control group ($M = 6:37$ AM). However, time of awakening was not associated with the main outcome variable, the PIPR.

Pupil diameter in response to the blue and red stimuli was plotted against time for representative SAD and control group participants in Figure 1. Means and standard deviations for the 7-second baseline pupil diameter, 30-second sustained response after light offset, PIPR (mm), and PIPR Change (%), Net PIPR, and Net PIPR Change (%) are shown in Table 2. Diagnostic groups did not differ on the measure of baseline pupil diameter prior to the first test stimulus ($F_{1,29} = 0.09$, ns), or on the measure of pupil diameter during the blue stimulus duration ($F_{1,29} = 1.46$, ns) or red stimulus duration ($F_{1,29} = 2.16$, ns). There was no relationship between baseline pupil diameter and net PIPR change ($r^2 = -0.16$, ns).

3.1 Diagnostic group comparisons

The control group showed a significantly larger PIPR in response to the blue stimulus than the SAD group ($F_{1,29} = 4.28$, $P < 0.05$), when controlling for age and gender (Table 2). Similarly, the percent change from baseline in response to the blue stimulus was significantly higher in controls compared to SAD participants ($F_{1,29} = 4.93$, $P < 0.05$). Groups did not differ on other pupil metrics (Table 2). Although means were in the hypothesized direction for net PIPR percent change, this difference was not statistically significant ($P = 0.12$). This is the first piece of evidence suggesting a diminished melanopsin-specific sensitivity in SAD.

3.2 Genotype comparisons

The mean and standard deviation for pupil values by genotype are shown in Table 3, along with genotype frequency by group. The blue PIPR was significantly different across genotypes of I394T, as the rare CC genotype had the largest blue PIPR (0.54 mm), the CT was intermediate (0.16 mm), and the TT genotype was lowest (-0.03 mm). Specific contrasts indicated that the TT and CT groups were significantly different from one another. When I394T C allele carriers (CC and CT genotypes; $n = 19$) were compared to non-carriers (TT genotypes; $n = 11$), the C allele carriers had a significantly higher mean blue PIPR compared to those without the C allele ($F_{1,29} = 6.21$, $P < 0.05$). We can only speculate that genetic variants may affect melanopsin sensitivity given that our sample size is small for typical genetic association tests.

4. Discussion

Our main study hypothesis, that the PIPR is diminished in SAD, was supported by the finding that the SAD group had a significantly smaller magnitude PIPR in response to the blue test stimulus compared to the control group. In addition, the PIPR percent change in response to the blue stimulus was significantly higher in the control group compared to the SAD group. However, the net PIPR and the net PIPR percent change were not significantly different, although group means were in the predicted direction. These results are unlikely to

be due to differences in retinal irradiance or melanopsin sensitivity as diagnostic groups and outcome variables did not vary based on variables expected to influence retinal irradiance including age, baseline pupil diameter, pupil diameter during the stimulus presentation, or retinal irradiance calculated post-hoc based on age and pupil diameter during the stimulus. In addition, diagnostic groups and outcome variables did not vary by factors that could affect the sensitivity of the melanopsin system such as time of assessment, photoperiod on the day of testing, and chronotype.

The I394T genotype was associated with the blue PIPR, but the P10L genotype was not associated with pupil measures. Because only 2 individuals had the rare I394T (CC) genotype and 3 individuals had the rare P10L (TT) genotype, these analyses are preliminary and likely underpowered. Others recently reported that I394T genotype interacts with stimulus intensity to predict the steady-state pupil diameter in healthy college students (Higuchi et al., 2013), which may be consistent with our findings here. However, Higuchi et al. (2003) did not measure the PIPR but rather measured steady state pupil diameter during stimulus presentation at various intensities, and did not genotype P10L, the variation found to be previously associated with SAD (Roecklein et al., 2009).

Both group means for chronotype fell in the intermediate range, although the SAD group mean trended towards relative eveningness compared to the control group. This is consistent with misalignment between the circadian clock and sleep/wake cycle seen in SAD (Lewy, 2007; Lewy et al., 2006). Time of awakening was consistent in that the SAD group reported waking later than controls. However, neither chronotype nor time of awakening were associated with the PIPR, so these variables are unlikely to explain the group difference found here on the PIPR.

To our knowledge this is the first study to test and report a diminished ipRGC-mediated blue PIPR in patients with SAD compared to controls with no history of depression. The implications and next steps for this line of research are detailed below.

4.1 Limitations

The magnitude of the PIPR to blue light observed in our study is lower (i.e., .01 mm in SAD and .23 mm in controls) than the values observed by others (e.g., .7 mm in controls; Gamlin et al., 2007). Although we used a light source of 13.7 log photons/cm²/s at the cornea, it was not presented in Maxwellian view, but rather was a diffuse light source presented without pharmacological pupil dilation. The calculated effective retinal irradiance was 12.3–12.5 log photons/cm²/s due to age dependent light absorption by the lens and due to pupil constriction during stimulus presentation. However, age, minimum pupil diameter during the stimulus, and calculated retinal irradiance were not associated with the magnitude of the blue PIPR, nor were those variables different between diagnostic groups, indicating that individual differences in retinal irradiance are unlikely to explain our findings. Because a threshold retinal irradiance of at least 12 log photons/cm²/s is required to produce the PIPR (Gamlin et al., 2007), our calculated retinal irradiance of between 12.3 and 12.5 log photons/cm²/s was on the very low end of irradiance needed to elicit the PIPR. Using higher irradiance, and either pupil dilation or a stimulus presented in Maxwellian view would further increase and standardize retinal irradiance across participants in future research. Future studies could also adjust stimuli before testing based on age-predicted lens absorption to better equate retinal irradiance across participants (Coren, 1987; Pokorny et al., 1987; Gamlin et al., 2007), although the standard deviations between calculated retinal irradiances in our study was only 0.19 log photons/cm²/s for the blue stimulus, and 0.16 log photons/cm²/s for the red stimulus.

An additional limitation may result from the current study's measurement of circadian time utilizing the MEQ, which is a strong but imperfect predictor of objective measures of circadian time such as dim-light melatonin onset (DLMO; Bernert et al., 2006). However, exact identification of circadian time may not be crucial for the current sample. Data from Zele et al. (2011) indicate that the magnitude of the PIPR is highest and relatively stable between CT3 and CT11, and in the evening the amplitude of the PIPR decreases about 2.4 hours prior to DLMO, an hour or so prior to bedtime. Therefore, since our participants were all studied during the predicted afternoon plateau of the maximum PIPR amplitude, and prior to the period of decrease in PIPR, it is unlikely that significant variation due to the timing of assessment or circadian phase is present in our study. However, others have found a similar peak PIPR during this time, but not a plateau (Munch et al., 2012). Therefore, future studies could measure DLMO to confirm that any effects of circadian variation on the PIPR are removed or controlled in group comparisons, ensuring that the results here, indicating that the diagnostic group difference is not due to circadian time, are indeed valid.

Recent light history can alter the sensitivity of melanopsin containing cells (Wong, et al., 2005), which may have an unknown effect on our measures. Importantly, circadian responses are more sensitive to light during short photoperiods when tested in Syrian hamsters (Glickman et al., 2012). Determining how photoperiod alters circadian responses to light (Glickman et al., 2012), including the effect on the PIPR, will be an important step for future studies with seasonal comparisons between human subjects. In our study, we attempted to control for light history by including photoperiod on the day of assessment as a covariate. No variance in dependent variables was accounted for by photoperiod and groups did not differ based on the photoperiod on the day of testing. A more accurate measurement of light received by a given individual prior to pupillometry could be collected in future studies with the use of light-sensitive actigraphy.

Subjects were tested with freely reactive pupils, while not under midriasis. A freely reactive pupil initially constricts in response to a stimulus, decreasing the light reaching the retina, leading to redilation until a steady state is achieved under constant illumination. The pupil diameter at steady state differs between individuals, therefore allowing differing amounts of light to reach the retina. Although differing steady state diameter could have affected the current results, pupil diameter and retinal irradiance based on pupil diameter were not associated with outcome variables or group membership in the present study and are therefore unlikely to have confounded our tests. Further, individuals under pharmacologically induced midriasis may also have different overall pupil sizes, and would also therefore differ in the amounts of light at the retina unless a Maxwellian presentation is used. Our use of freely reactive pupils and non-Maxwellian presentation are more similar to the naturalistic responses of this circadian photoreceptive system in humans. In addition, other authors have presented stimuli unilaterally, and measured the consensual response in the non-dominant eye to control for differences in retinal irradiance. In the present case, stimuli were presented bilaterally, and measured in the left eye while dominance was not assessed. It is likely that bilateral presentation caused the diameter during stimulus presentation to be smaller than it would be with midriasis, thereby reducing the retinal irradiance. However, retinal irradiance was calculated post-hoc, taking pupil diameter during stimulus presentation, and age, into account. Although this may introduce noise into measurement, lack of midriasis is a more naturalistic measurement setting. In addition, the comparison here of red and blue light is confounded by the difference in retinal irradiance due to selective lens absorption of blue wavelengths. In future studies, the retinal irradiance should be equated (ours were 12.3 blue vs. 12.3 red log photons/cm²/s) allowing for a comparison of the red and blue responses.

The age of participants ranged from 20 to 60 years, and was relatively evenly distributed across this range, with no evidence of skewness or kurtosis (e.g., there were not more older than younger participants). However, lens density increases with increasing age. In the present study, we did not have the ability to increase the stimulus intensity for older individuals to ensure equal retinal irradiance across participants, but age and gender were covaried in our analyses, and age-dependent retinal irradiance was calculated and found to not differ between diagnostic groups. Since age is evenly distributed across old and younger individuals in this study, and participants were matched by age, the effect of age on retinal irradiance is not expected to explain the group difference found in this study.

Nine participants in the SAD group, and two participants in the control group reported taking medications or legal drugs that could potentially affect pupil diameter. Pupil responsiveness can be affected by drugs that affect both the sympathetic (e.g., nicotine, bupropion) and parasympathetic (e.g., caffeine, clomipramine) autonomic nervous system either agonistically or antagonistically. In the present study, participants were not excluded on the basis of medications, but this could be done in future studies to further control the potential effects of these medications and prevent any potential group differences. Although more individuals in the SAD group were taking medications compared to the control group, post hoc analyses confirmed that medication status was not associated with PIPR magnitude (data not shown). Including individuals taking medications increases the external validity of our results, but medication use will continue to be measured and controlled for in future studies.

The retinal subsensitivity hypothesis of SAD proposes that a seasonal variation in retinal sensitivity, with winter subsensitivity and normal functioning in summer, may explain SAD. Future studies should measure the PIPR in SAD and control participants in both summer and winter to determine if one of two situations exist: (1) retinal sensitivity may not increase in winter in SAD as predicted by the retinal subsensitivity hypothesis, or (2) individuals with SAD have lower melanopsin-specific sensitivity all year, which is only problematic when winter light levels fall below a given threshold.

Finally, we did not record the refraction or prescription for participants with corrected vision, or covary for whether or not participants wore contact lenses, although we did not have any participants who used glasses. Because some contact lenses are tinted, and tinting could cause attenuation of specific wavelengths of light, this will be recorded in future studies.

4.2 Conclusions

The present study indicates that the melanopsin-specific PIPR is less pronounced in SAD during winter compared to controls. If the reduced PIPR is confirmed to be due to SAD group membership rather than diabetes, glaucoma, retinitis pigmentosa or other systemic diseases, then the PIPR may represent a biomarker of SAD. In the present study, participants in both groups reported being free of diabetes, diabetic retinopathy, glaucoma and retinitis pigmentosa, but these and other systemic diseases will need to be more rigorously assessed in future studies. If retinal subsensitivity in SAD is mediated at least in part by reduced melanopsin sensitivity, a few implications are possible. Given the role of melanopsin in both circadian and acute behavioral effects of light, it is possible that reduced melanopsin sensitivity would lead to impaired photoentrainment of the circadian clock, reduced alertness, or even changes in sleep homeostasis, all of which may be etiologically significant in SAD (Stephenson et al., 2011). For example, men with SAD have a longer duration of melatonin release in winter compared to summer (Wehr et al., 2001). In humans, short wavelength light is the most effective for acute suppression of melatonin, indicating a primary role for melanopsin in melatonin suppression (Brainard et al., 2001; Thapan et al.,

2001; Revell et al., 2010). Reduced melanopsin sensitivity, if confirmed upon replication, is likely to be only one of multiple risk factors in SAD, with others including other photoreceptors, or serotonergically mediated reduced sensitivity of the central clock to light (Stephenson et al., 2011).

In the future, it will be important to evaluate stimulus irradiance levels and threshold effects, control for recent light history and circadian time, and then determine the clinical significance of the PIPR if these results are replicated. Clinical challenges remaining in SAD treatment include that only about 53% of individuals with SAD respond to light therapy (Terman et al., 1989) and only 41% of those adhere to the treatment in future winters (Schwartz et al., 1996), possibly because the treatment as prescribed is not fully effective. Treatment response can also be incomplete when compared to summer spontaneous remission (Postolache et al., 1998). Further understanding of melanopsin's etiological impact on SAD may improve treatment. If ipRGCs are hypofunctional in some or even most individuals with SAD, two possibilities exist: (1) light therapy may not work at all for these individuals, or (2) light therapy may need to be a higher intensity or longer duration than for other individuals with SAD. Until these two options are explored empirically, the mechanism behind melanopsins' potential role in light therapy would be speculative. Studies have established that blue or blue-enriched lights at significantly lower light irradiances or durations, yield similar rates of improvement in SAD as traditional bright white light therapy units (Glickman et al., 2006; Anderson et al., 2009; Meesters et al., 2011), and are able to phase advance rhythms (Smith et al., 2009). Outstanding questions include whether the melanopsin cells are primary in mediating light therapy's effects, and whether it may be possible to potentiate the effects of light therapy by using long wavelength light first (Rollag, 2008; Mure et al., 2009; Gooley et al., 2010). Furthermore, it is possible that the PIPR could be used to predict the timing, duration, and potential benefit of light therapy, opening the door to individualized treatment prescriptions in SAD. For example, given that some individuals with SAD don't respond to light therapy, the PIPR might help predict which individuals should try antidepressants (e.g., Moscovitch et al., 2004; Modell et al., 2005; Lam et al., 2006) or cognitive behavioral therapy for SAD (Rohan et al., 2004; 2007; 2009). It could be, however, that subgroups of individuals with a diminished PIPR could change the light therapy time of administration, spectral composition, light intensity, or duration to achieve better results. Our work is preliminary, based on the limitations listed above, but this line of research has the potential to identify a simple and non-invasive neurological test for psychopathology.

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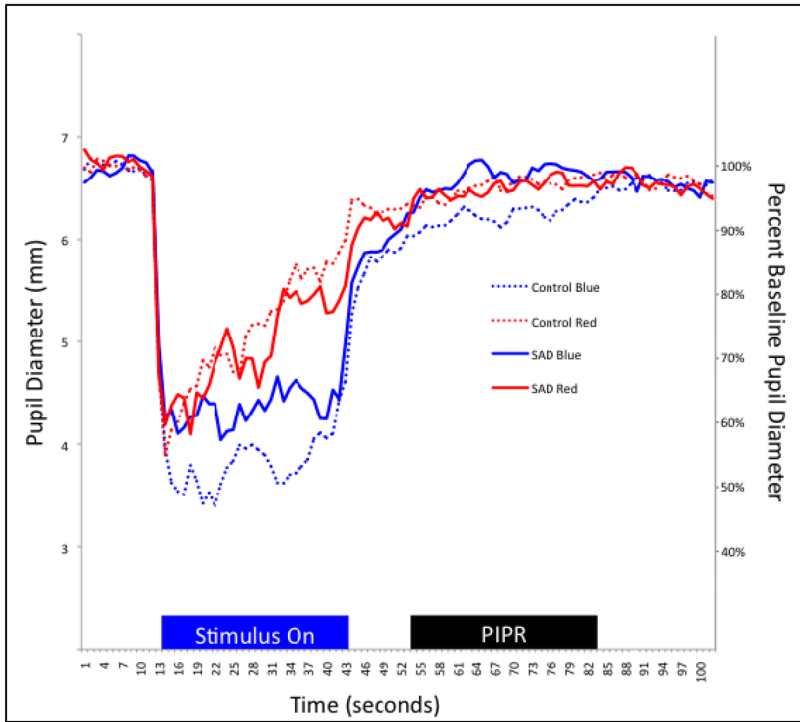


Figure 1. Time trace plot of the pupillary response to the blue and red stimuli for one representative SAD participant and one Control participant. Notes: Stimulus duration, shown in the blue “Stimulus On” block, was for 30 seconds, following 7 seconds of baseline recording. PIPR, shown in the black “PIPR” block, was measured starting 10 seconds after light offset for 30 seconds. Oscillations evident in the individual traces are attributed to the lack of midriasis and stimulus presentation to both eyes, including the eye being measured.

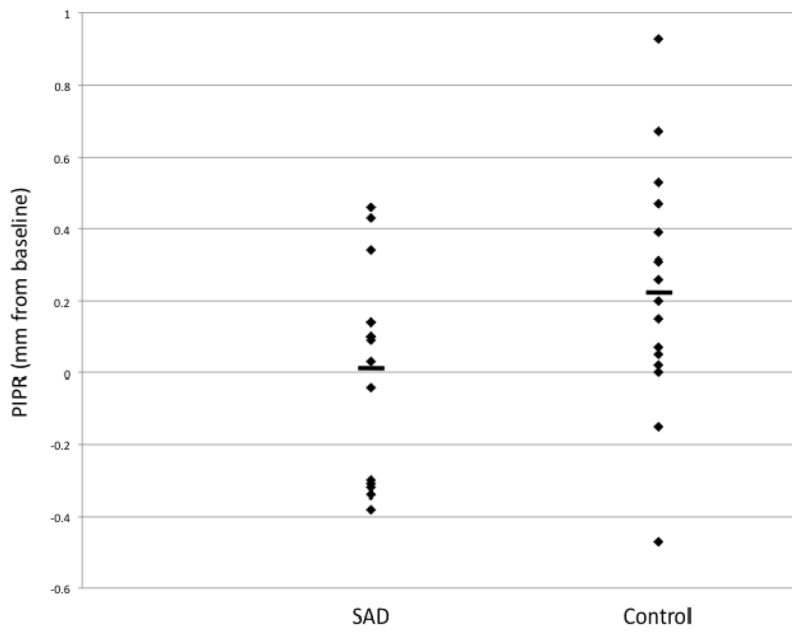


Figure 2. Post-Illumination Pupil Response (PIPR) values expressed in mm as change from baseline pupil diameter, between SAD and control groups.
 Note: The mean post-illumination pupil response (PIPR) is marked with a horizontal bar for each group. PIPR was measured from 10–40 seconds after stimulus off, and is presented here as a change from baseline. Baseline was calculated for the seven seconds prior to stimulus on.

Table 1

Demographic Characteristics and Baseline Measures Compared Between SAD Participants and Controls; $M(SD)$.

	SAD (<i>n</i> = 15)	Controls (<i>n</i> = 15)
Age (y)	36.1(14.8)	37.4(14.7)
% Female	86.8%	73.4%
Baseline pupil diameter (mm)	5.73(1.14)	5.99(1.03)
MEQ ^a	48.4(12.8)	54.5(8.8)
Daylength on assessment (hh:mm)	11:03(0:34)	11:18(0:33)
Time of assessment (hh:mm)	15:33(1:06)	15:31(1:37)
Calculated retinal irradiance		
Blue (log photons/cm ² /s)	12.27(.22)	12.25(.18)
Red (log photons/cm ² /s)	12.49(.18)	12.44(.14)

All *P*s were greater than 0.05 (*ns*).

^aMEQ - Morningness Eveningness Questionnaire measure of chronotype.

Table 2

Diagnostic group comparisons on PIPR outcome variables.

	SAD			Controls		
	R	B	B-R (Net)	R	B	B-R (Net)
Baseline pupil (mm)	5.59(1.15)	5.73(1.14)	0.15(0.35)	5.80(1.14)	5.99(1.03)	0.19(0.40)
Stimulus (mm)	4.01(1.06)	3.99(1.04)	-0.03(0.31)	3.79(0.72)	3.79(0.60)	-0.01(0.25)
Sustained (mm)	5.65(1.05)	5.86(1.05)	0.21(0.29)	5.78(0.96)	5.76(1.05)	-0.02(0.32)
PIPR (mm)	-0.063(0.68)	0.01(0.28)*	-0.07(0.27)	0.02(0.47)	0.23(0.34)*	0.21(0.60)
PIPR Change (%)	-2.29(12.3)	-0.79(5.56)*	-0.86(5.73)	-0.50(8.72)	3.85(5.88)*	4.35(11.22)

Mean values are shown in mm or percentages, followed by the Standard Deviation in parentheses; Baseline - Average pupil diameter 7 seconds prior to light onset; Stimulus - Average diameter over 30 seconds during stimulus on; Sustained - Average diameter over 30 seconds, starting 10 seconds after light offset; B - blue stimulus; R - red stimulus; B-R - Net difference between values of either PIPR or PIPR percent change for the blue and red stimuli.

* $P < 0.05$

Table 3

Melanopsin genotype groups comparisons; *M*(*SD*) or *n*(%).

	P10L			I394T			P
	CC	CT	TT	CC	CT	TT	
<i>N</i>	17	9	3	2	17	11	
Baseline pupil (mm)	5.65(1.04)	6.16(1.33)	6.14(0.42)	5.75(0.09)	5.92(1.24)	5.78(0.94)	<i>ns</i>
blue PIPR (mm)	0.12(0.34)	0.17(0.33)	-0.10(0.32)	0.54(0.56)	0.16(0.30)	-0.03(0.27)	<0.05
red PIPR (mm)	-0.11(0.64)	0.22(0.50)	-0.20(0.32)	-0.31(0.94)	0.04(0.65)	-0.08(0.40)	<i>ns</i>
Net PIPR (mm)	-0.01(0.60)	0.00(0.57)	1.00(1.39)	0.43(1.10)	-0.05(0.55)	0.24(0.91)	<i>ns</i>
Net PIPR Change (%)	2.37(11.4)	0.03(5.28)	2.17(5.26)	17.17(29.34)	-0.05(7.01)	1.71(5.02)	<i>ns</i>
SAD group	8(50%)	6(38%)	2(12%)	8(50%)	7(44%)	1(6%)	<i>ns</i>
Control group	10(71.5%)	3(21.5%)	1(7%)	3(20%)	11(73%)	1(7%)	

Notes: *ns* > 0.05; P10L – rs2675703; I394T – rs107961