1 RESEARCH ARTICLE

2 RUNNING HEAD: Eye closure impacts visual processing

³ Differential cortical and subcortical visual processing with

4 eyes shut

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18 ABSTRACT

19 Closing our eyes largely shuts down our ability to see. That said, our eyelids still pass some light, 20 allowing our visual system to coarsely process information about visual scenes, such as changes in 21 luminance. However, the specific impact of eye closure on processing within the early visual system 22 remains largely unknown. To understand how visual processing is modulated when eyes are shut, we 23 used functional magnetic resonance imaging (fMRI) to measure responses to a flickering visual stimulus 24 at high (100%) and low (10%) temporal contrasts, while participants viewed the stimuli with their eyes 25 open or closed. Interestingly, we discovered that eye closure produced a qualitatively distinct pattern of 26 effects across the visual thalamus and visual cortex. We found that with eyes open, low temporal 27 contrast stimuli produced smaller responses, across the lateral geniculate nucleus (LGN), primary (V1) 28 and extrastriate visual cortex (V2). However, with eyes closed, we discovered that the LGN and V1 29 maintained similar BOLD responses as the eyes open condition, despite the suppressed visual input 30 through the eyelid. In contrast, V2 and V3 had strongly attenuated BOLD response when eyes were 31 closed, regardless of temporal contrast. Our findings reveal a qualitatively distinct pattern of visual

- 32 processing when the eyes are closed one that is not simply an overall attenuation, but rather reflects
- distinct responses across visual thalamocortical networks, wherein the earliest stages of processing
- 34 preserves information about stimuli but is then gated off downstream in visual cortex.

35 NEW & NOTEWORTHY

- 36 When we close our eyes coarse luminance information is still accessible by the visual system. Using
- 37 functional magnetic resonance imaging, we examined whether eyelid closure plays a unique role in
- visual processing. We discovered that while the LGN and V1 show equivalent responses when the eyes

- 39 are open or closed, extrastriate cortex exhibited attenuated responses with eye closure. This suggests
- 40 that when the eyes are closed, downstream visual processing is blind to this information.
- 41 Keywords: luminance, eye closure, fMRI, LGN, visual cortex
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43 **INTRODUCTION**

44 Light exposure during sleep has substantial effects on the brain: it can alter circadian rhythms, 45 sleep quality, and mood (1,2). During sleep, our eyes are closed and the eyelids function as potent filters of visual information. However, our eyelids are only partial filters and do not completely attenuate all 46 47 visual information (3,4). The eyelid has been characterized as a red-pass filter, with an estimated 6% red 48 light spectral transmittance (3). Indeed, subjective experience with high luminance stimuli, such as 49 during a sunny day, corroborates the idea that changes in luminance are still detectable when our eyes are closed. With partial, rather than complete, filtering properties, it follows that the visual system 50 51 processes external visual information with our eyes closed, as well.

52 How does the visual system process information when our eyes are closed? It is possible that 53 the filtering properties of the eyelid simply quantitatively suppress responses across visual regions, due 54 to the attenuation of input. Alternatively, eye closure could induce qualitatively distinct changes in 55 visual response, selectively modulating responses in specific brain networks. While little is known about 56 stimulus-evoked visual responses with eyes closed, resting-state fMRI studies have investigated 57 spontaneous dynamics during eye closure in the absence of any visual stimulus presentation (5-7). These 58 studies found differences in resting-state functional connectivity in attentional networks depending on 59 whether eyes were open or closed, along with differences in activation in prefrontal cortex, parietal and 60 frontal eye fields, and LGN. While eye closure appears to play a unique role in modulating brain 61 responses, the impact that eye closure has on stimulus-evoked visual responses remains poorly 62 understood.

63 In this study, we sought to shed light on the role that eye closure plays in modulating responses 64 within the visual processing hierarchy. To do so, we measured fMRI BOLD responses within visual cortex 65 and subcortex while participants viewed high and low intensity visual stimuli, with their eyes open or 66 shut. We manipulated the intensity of visual input via temporal contrast modulation, in which the 67 luminance of a uniform visual stimuli flickered rapidly between extreme whites and blacks (high 68 temporal contrast), or between middling intensities (low temporal contrast). Indeed, previous work has 69 shown visuocortical responses to be sensitive to changes in luminance (8). By measuring BOLD 70 responses to high and low luminance contrast stimuli, we examined whether there is a qualitatively 71 unique pattern of luminance responses across the visuocortical hierarchy when one's eyes are closed, 72 compared to when they are open.

73 MATERIALS AND METHODS

74 Participants

Data was acquired from a total of 8 healthy participants (5 females, 3 males; 3 Asian, 1 Black or African American; 4 White). Participants were aged 18-35 years, reported normal or corrected-tonormal visual acuity, and were recruited from Boston University and the surrounding community. All participants provided written informed consent before study enrollment and completed a metal screening form indicating that they had no MRI contraindications. Participants were reimbursed for their study participation.

81 Apparatus & stimuli

82 Stimuli were generated using custom software written in MATLAB (version 2019b) in 83 conjunction with Psychtoolbox (9). Participants viewed stimuli that was back-projected onto a screen set within the MRI scanner, using a ProPIXX DLP LED (VPixx Technologies) projector system (minimum 84 luminance: 1.2 cd/m2; maximum luminance: 2507.9 cd/m²). Photometer measurements (model LS-100; 85 86 Konica Minolta) carried out before the study were used to verify the linearity of the display (1 digital-to-87 analog conversion (DAC) step = 9.835 cd/m^2). These measurements were used to calculate the stimulus 88 luminance and were acquired from the inner-facing side of the back-projection screen while positioned within the MRI scanner bore. This was done to best account for the attenuation in luminance due to 89 90 back-projection screen characteristics.

91 During each functional run, participants fixated on a median luminance crosshair at the center 92 of the display while shown a full screen flickering display (17 degrees of visual angle) with no spatial 93 contrast (Figure 1). The full field flicker was presented in a block design with three trial types (baseline, 94 high, and low temporal luminance contrast), with each event lasting 16 seconds. In the baseline events, 95 the full field display was a constant median luminance with no luminance modulation. During high 96 events, the full field display flickered with an amplitude envelope of 100% around the middle luminance 97 value. For low events the full field display flickered with an amplitude envelope of 10% around the 98 median luminance value. All high and low events flickered at a frequency of 7 Hz.

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102 Experimental design

Subjects participated in two scan sessions, each lasting approximately two hours. The first session was dedicated to collecting anatomical images and data for population receptive field (pRF) mapping using standard techniques and stimuli (10,11). The second session was dedicated to collecting proton-density (PD) weighted anatomical imaging and fMRI blood oxygenation level-dependent (BOLD) data across the eyes open and closed conditions, during the luminance task.

108 During the second experimental session, we collected three PD-weighted anatomical scans. PD-109 weighted anatomical imaging has previously been used to better localize the LGN (12,13). Following the 110 PD-weighted scans, participants completed three consecutive runs of a functional localizer. The visual stimulus for the functional localizer contained a full field flickering grating stimulus (diameter = 6.0°) 111 112 with a centered circle (diameter = 0.8°). Within the centered circle, letters rapidly appeared one at a 113 time with a new letter appearing every 200 ms. Participants were instructed to press a button whenever 114 the letters 'J' and 'K' appeared within the centered circle. During the localizer blocks, the full field display alternated between a flickering grating stimulus and a full field non-flickering display at median 115 luminance value. Participants completed 12 total blocks (6 flickering field, 6 non-flickering field) with an 116

extra non-flickering block at the beginning of the run. At the end of each localizer run, participants wereasked to report their wakefulness level.

119 Participants then completed the luminance flicker task. The task began and ended with a baseline event. High and low temporal contrast conditions were pseudo-randomly ordered, with all high 120 121 and low events interleaved with a baseline event. Each run contained 12 events (6 high, 6 low) 122 interspersed with 12 baseline events, lasting a total of 384 seconds. On each run participants were 123 instructed to press a button after each full breath cycle (1 inhale, 1 exhale). This button task was chosen 124 to ensure that participants did not fall asleep and engaged with the task, while not requiring eyes to be 125 open. For each run, participants were instructed to either keep their eyes open and fixate on the 126 crosshair or to keep their eyes closed throughout the run. Each scan session began with an eyes-closed 127 run, and consecutive runs alternated between open and closed conditions. We always began with the 128 eyes closed condition to ensure we acquired a sufficient number of runs in this condition, where BOLD 129 modulations may be lower compared to eyes-open runs. To ensure participants kept their eyes closed or 130 open, real time eye monitoring was carried out using an EyeLink1000, for the duration of each run. On 131 average, we collected 5 runs with eyes closed and 4 runs with eyes open, for each subject.

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133 MRI data acquisition

134 All neuroimaging data were acquired using a research-dedicated Siemens Prisma 3T scanner 135 using a Siemens 64-channel head coil. A whole brain anatomical scan was acquired using a T1-weighted 136 multi-echo MPRAGE (1 mm isotropic voxels; field of view (FOV) = 192 x 192 x 134 mm, flip angle (FA) = 7.00°, repetition time (TR) = 2200 ms, echo time (TE) = 1.57 ms). Proton density (PD)-weighted 137 138 anatomical scans were acquired to localize LGN (0.9mm x 0.9mm x 1.7mm; TR = 2950.0 ms; TE = 15.6 139 ms; FA = 180°). Functional scans were acquired using T2*-weighted in-plane simultaneous imaging (2 140 mm isotropic voxels; FOV = $104 \times 104 \times 70$ mm, FA = 64.00° , TR = 1000 ms, TE = 30 ms, SMS factor = 5, 141 GRAPPA acceleration = 2).

142 Anatomical data analysis

143 T1-weighted anatomical data were analyzed using the standard "recon-all" pipeline provided by 144 the FreeSurfer neuroimaging analysis package (14), generating cortical surface models, whole brain 145 segmentations, and cortical parcellations. All PD-weighted scans were aligned to each subject's 146 anatomical space and averaged together (using AFNI's 3dcalc).

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148 Functional data analysis

149 Functional BOLD time-series data were first corrected for echo-planar imaging (EPI) distortions 150 using a reverse phase-encode method implemented in FSL (15) and were then preprocessed with FS-FAST using standard motion-correction procedures, slice timing correction, and boundary-based 151 registration between functional and anatomical spaces (16). To optimize spatial precision of 152 experimental data, no volumetric spatial smoothing was performed (full-width half-maximum 0 mm). To 153 154 achieve precise alignment of experimental data within the session, cross-run within-modality robust 155 rigid registration was performed, using the middle time point of each run (17). BOLD time-series data 156 were demeaned and converted to units of percent signal change. Data collected during the separate pRF 157 mapping scans were analyzed using the analyzePRF toolbox (11). Results from the pRF model were used 158 to manually draw labels for our regions of interest within visual cortex.

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160 Statistical analysis

161 The results from the pRF modeling were used to identify region-of-interest (ROI) labels for each cortical region before analysis. ROI labels included voxels located inside the cortical ribbon for 162 V1/V2/V3, which were identified using a visual area network label generated using an intrinsic 163 164 functional connectivity atlas (18). Results from the pRF modeling were additionally used to select voxels 165 with visual field eccentricity preferences less than 17 degrees visual angle away from fixation as this was 166 the measured extent of the screen within the MRI scanner. Cortical voxels with a poor pRF model fit ($r^2 < r^2$ 0.10) were removed from further analyses. Initial LGN labels were acquired from thalamic segmentation 167 168 and parcellation in Free-Surfer for each participant. These initial labels were overlaid with the GLM 169 results from the functional localizer and the PD-weighted scans, and only intersecting voxels were 170 chosen for the final LGN labels and further analyses.

An event-triggered average was computed for each flickering condition (low and high) per eyelid 171 172 condition and ROI. The BOLD time-series for each ROI per run was separated by the low and high trials, 173 and all trials of a given type were averaged together. Average BOLD magnitude in response to the 174 stimulus presentation was computed by averaging 4-16 s post-stimulus onset for each trial. Two-way 175 between-subjects ANOVA were performed to test for any main effects of temporal contrast and eye 176 closure and any interaction of the two on average BOLD magnitude during stimulus presentation. 177 Additional event-triggered average analysis was done with eccentricity, in which the time-series for V1/V2/V3 voxels were first separated into eccentricity bins defined by degree visual angle relative to 178 179 fixation. Foveal-tuned voxels were between $0.01^{\circ} - 1.5^{\circ}$, parafoveal-tuned voxels were between $1.5^{\circ} - 1.5^{\circ}$ 4.0° , and peripheral-tuned voxels were between $4.0^{\circ} - 17.0^{\circ}$. An additional ANOVA was performed to 180 test for any main effect of eccentricity on BOLD response during stimulus presentation. Multiple 181 182 comparison correction was done using Bonferroni correction of α/n at a familywise α of 0.05 where n is 183 the number of tests performed.

184

185 **RESULTS**

186 We first examined how temporal contrast modulated thalamic and visuocortical responses, and 187 if eye closure impacted these responses. With eyes open, LGN, V1, and V2 showed larger responses to 188 high temporal contrast stimuli, compared to low temporal contrast stimuli (Figure 2A). Indeed, during 189 eyes open with high temporal contrast stimuli, all ROIs had significantly elevated BOLD responses [LGN: t(7) = 3.27, P = 0.006; V1: t(7) = 5.16, P < 0.0001; V2: t(7) = 3.40, P = 0.005; V3: t(7) = 2.54, P = 0.019),190 191 though the significant response in V3 did not survive multiple comparisons correction. When the 192 participants closed their eyes, however, LGN and V1 maintained their stronger responses to higher 193 contrast stimuli [LGN: F(1,31) = 4.31, P = 0.047; V1: F(1,31) = 11.05, P = 0.002], which did not differ from their eyes closed conditions [LGN: F(1,31) = 0.02, P = 0.975; V1: F(1,31) = 1.74, P = 0.20]. In other words, 194 195 while responses in LGN and V1 were significantly modulated by temporal contrast, they were 196 completely unaffected by eye closure, despite the profound suppression of visual input from the eyelid.

197 Interestingly, while eye closure did not appear to have a major effect on the earliest stages of 198 visual processing (LGN and V1), we observed a qualitatively distinct pattern within extrastriate cortices 199 V2 and V3. When the eyes were closed, there was a drastic attenuation of stimulus evoked responses, 200 regardless of temporal contrast [Main effects of eye closure: V2: F(1,31) = 6.45, P = 0.017; V3: F(1,31) =201 5.79, P = 0.02; Main effect of temporal contrast V2: F(1,31) = 2.91, P = 0.09; V3: F(1,31) = 0.54, P = 0.47]. 202 Pairwise comparisons revealed a significant decrease in BOLD response to high temporal contrast stimuli 203 with eye closure in V2 (t(14)=-3.13; P = 0.003) and V3 (t(14)=-3.09; P = 0.003). Overall, these results 204 indicate that visual processing appears to be qualitatively different with eyes closed compared to when 205 eyes are open. The BOLD response in LGN and V1 was modulated by temporal contrast but was

unaffected by eye closure, whereas eye closure strongly reduced responses in extrastriate cortices V2and V3.



²⁰⁸ Figure 2. 209

210 Along with the heterogeneity in patterns observed across striate and extrastriate regions, it is 211 possible that there exists heterogeneity within each region. It has been reported that there is an 212 eccentricity bias of the BOLD response in V1 and V2, when participants viewed center-surround stimuli with no local contrast (19). To test for an eccentricity bias and if eye closure impacts this bias, we 213 214 separated voxels in V1-V3 by their eccentricity preference, based on pRF estimates (LGN was excluded 215 from this analysis due to being underpowered for pRF analyses). We defined foveally-preferring voxels 216 as those preferring between $0.01^{\circ} - 1.5^{\circ}$ from fixation, parafoveal-preferring voxels were those between $1.5^{\circ} - 4.0^{\circ}$, and peripheral-preferring voxels were between $4.0^{\circ} - 17.0^{\circ}$. As low temporal 217 218 contrast trials elicited no significant activation across visuocortical regions, we did not test for an effect 219 of eccentricity during low temporal contrast trials. We found that the effect of eccentricity was not 220 significant in V1 [F(2,47) = 1.16, P = 0.333] (Figure 3), nor in V2 [F(2,47) = 0.51, P = 0.606] nor V3 [F(2,47)221 = 0.30, P = 0.744]. No ROIs had any significant interaction between eye closure and eccentricity [V1: F(2,47) = 0.26, P = 0.772; V2: F(2,47) = 0.94, P = 0.397; V3: F(2,47) = 0.26, P = 0.768]. This suggests that 222 223 across striate and extrastriate cortices there is no eccentricity bias in BOLD responses nor any difference with eye closure. Thus, the impact of high temporal contrast stimuli and eye closure on BOLD appear 224 225 uniform within each visuocortical area.



226 Figure 3.

227 **DISCUSSION**

With subjective experience it is clear that we can still perceive visual stimuli with closed eyes, but how distinct stages of the visual system supported this filtered visual experience was unknown. In this study, we found that eye closure produces a qualitatively distinct pattern of modulatory responses within the early visual system: closing one's eyes selectively attenuated luminance processing in extrastriate cortex, but not in LGN nor striate cortex.

233 In line with previous literature showing that early visual responses can still occur when the eyes 234 are closed (5,20), we demonstrated that with closed eyes, luminance-dependent responses remain 235 present in the LGN and V1. However, we found substantial heterogeneity in activation across regions 236 when eyes were closed. One hypothesis as to why we observed strongly attenuated BOLD with closed 237 eyes in extrastriate cortex, but not the LGN nor striate cortex, is that top-down modulation of 238 visuocortical responses is often stronger in extrastriate compared to striate cortex (21-23). It has been 239 demonstrated that higher-order sensory regions, such as the frontal eye field (FEF), may account for the 240 selective top-down modulation of extrastriate cortical responses (24). Resting-state fMRI studies that 241 examined altered functional connectivity between eyes open and closed states found increased 242 activation of the FEF during eyes closed relative to eyes open scans (7), lending further support to top-243 down modulation of extrastriate cortex during eyes closed states. Interestingly, one study which 244 microstimulated the FEF of monkeys and measured visuocortical responses with fMRI found that FEF 245 stimulation modulated extrastriate areas only in the presence of a visual stimulus, indicating that top-246 down modulation of the extrastriate cortices is dependent on bottom-up influence (25). Since our 247 paradigm includes a visual stimulus, it is possible that eye closure in the presence of visual stimuli 248 attenuates extrastriate cortical responses through both top-down and bottom-up mechanisms. Additionally, the eyelid abolishes almost all structure and form-like information, which is necessary to 249 250 elicit responses in extrastriate cortices that prefer higher-level feature selectivity, such as spatial 251 contrast, shapes, and contours. However, eyelid closure still passes through luminance information,

which is known to activate striate cortex (8). This preservation of luminance information, but attenuation of higher-level information, may explain the preservation of early visual pathway activation with weakened extrastriate activation.

255 Visuocortical responses have been shown to depend on luminance modulation, with responses 256 increasing monotonically with luminance (8). In addition to luminance modulation, luminance response 257 functions are strongly contrast dependent, with lower spatial contrast drastically decreasing 258 visuocortical responses to luminance (8). Since the eyelid filters out much visual information, it is likely 259 that spatial contrast no longer impacts visual responses and that luminance information dominates what 260 might pass through the eyelid. Additionally, the lower luminance retinal input with eye closure cannot 261 fully explain our results since LGN and V1 showed no significant change in BOLD activation between 262 open and closed eye conditions. Since the eyelid is characterized as a red-pass filter (3), it is possible 263 that early visual pathways preferentially process this red visual content that extrastriate cortex is blind 264 to; however, to our knowledge no evidence of this exists. Although further work will be needed to 265 better unpack luminance responses in the early visual system, our results suggest that luminance-based 266 responses within early visual areas may not always necessitate the existence of spatial contrast in order 267 to reveal themselves, as previously suggested.

268 There did not appear to be any significant effects of eccentricity on luminance responses, 269 neither with eyes open nor eyes closed. Previous research found an eccentricity bias in BOLD responses 270 in V1 and V2 to center-surround stimuli with no local contrast, most strongly at the edge between the 271 center and surround of the stimuli (19). However, we may not have observed a similar effect simply 272 because our visual stimulus did not include any spatial contrast, thereby precluding any edge effects 273 from a center-surround stimulus. This suggests that the luminance-dependent effect of eccentricity 274 might depend on where in the visual field an edge exists, which is consistent with another study 275 examining luxotonic responses in the visual cortex using fMRI, which also did not find any effect of 276 eccentricity on luxotonicity (8).

277 Our stimulus was designed to provide similar features of visual input across conditions, by 278 presenting a diffuse light with no spatial contrast, but the input was nevertheless not identical when the 279 eyes were closed due to the additional attenuation from the eyelid. Future experiments could test a 280 stimulus that mimics the reduced retinal input of the eyelid. This would involve measuring the spectral 281 transmittance of each participant's eyelid and adjusting the high temporal contrast visual stimulus to 282 account for the attenuated transmittance. Such a stimulus would necessarily abolish all structure and 283 form uniformly across the visual scene, which under standard models of center-surround neuronal 284 receptive field organization would predict no net change in visuocortical responses due to equivalent 285 stimulation across excitatory and inhibitory components of the visual cortex (26,27). However, recent 286 findings of luminance modulation within the visual cortex suggests that luminance information alone 287 can drive the visual system (8). Thus, a stimulus mimicking the eyelid preserves luminance information 288 and would be expected to still activate the visual system, in accordance with our eyes closed condition 289 in which high intensity stimuli still activates LGN and V1. If the visual thalamus and visuocortical 290 responses between a stimulus mimicking the eyelid with open eyes and the high intensity stimulus in 291 our eyes closed condition did not align, this would suggest that the physical properties of the eyelid are 292 not sufficient to explain our results.

An alternative explanation of our results is that modulation of visual processing during eye closure may be dependent on brain state, not just the physical barrier of the eyelid. Eye closure likely induces a change in overall brain state that alters both the processing of visual information and largescale functional network processing. Eye closure decreases activity in attentional systems in the occipital and parietal lobes and increases functional coupling between sensory thalamus and somatosensory regions (5-7). Even during short eye blinks, there is evidence of increased activity in the default mode network (DMN) and decreased activity in the dorsal attention network, suggesting attentional 300 disengagement even with short eye closures (28). During longer eye closures, as examined in this study, 301 we might expect prolonged or even greater magnitude DMN activation, supporting this idea of 302 attentional disengagement with closed eyes. These differences in spontaneous brain activity across 303 sensory and attentional systems point to altered brain states with eye closure. Exteroceptive and 304 interoceptive mental state hypotheses have been formulated where an exteroceptive mental state is 305 characterized by increased attention and sensory processing of the external environment with eyes 306 open (5). On the other hand, an interoceptive mental state is characterized by internally-directed 307 cognition and reduced sensory processing with eyes closed. Many brain states require prolonged periods of eye closure, such as sleep and meditation, that involve reduced sensory awareness of 308 309 external stimuli and enhanced internally-directed attention. Thus, eye closure may modulate visual 310 processing through attentional or brain-state-dependent mechanisms.

311

312 DATA AVAILABILITY

313 Source data for this study are openly available at DOI: 10.18112/openneuro.ds005194.v1.1.0

314 IRB STATEMENT

All aspects of the study were approved by Boston University's Institutional Review Board.

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331 The authors declare no competing financial interests

332 **DISCLAIMERS**

333 The authors declare no disclaimers

334 AUTHOR CONTRIBUTIONS

- 335 Conceived and designed research: NGC, MK, LDL, SL
- 336 Performed experiments: NGC, MK
- 337 Analyzed data: NGC
- 338 Interpreted results of experiments: NGC, MK, LDL, SL
- 339 Prepared figures: NGC
- 340 Drafted manuscript: NGC
- 341 Edited and revised manuscript: NGC, MK, LDL, SL
- 342 Approved final version of manuscript: NGC, MK, LDL, SL
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 default mode network while viewing videos. *Proc Natl Acad Sci USA* 110: 702-706, 2013.

405

406 FIGURE LEGENDS

407 *Figure 1.* Experimental design with sample stimulus frames displaying the high temporal

408 **contrast and low temporal contrast displays.** High temporal contrast flickered at 7 Hz with a

- luminance amplitude envelope of 100%, encompassing the maximum (255 a.u.) and minimum (0 a.u.)
 possible luminance values. The low temporal contrast events also flickered at 7 Hz with a luminance
- amplitude envelope of 10%, encompassing a range of luminance values between 140 a.u. and 115a.u.
- 413

414 Figure 2. Eye closure has minimal effect on visual responses in LGN and V1, while suppressing 415 responses in V2 and V3. (A) Event-triggered average for luminance task across ROI and eye 416 condition. Across LGN, V1, and V2, during eyes open runs high temporal contrast stimuli elicits a 417 greater BOLD response than with low temporal contrast stimuli. Though there is no effect of temporal 418 contrast in V3, BOLD response increases regardless of the stimulus temporal contrast. During eye 419 closure, BOLD responses in LGN and V1 during the high temporal contrast stimuli elicits a similar 420 BOLD response as during eyes open runs. With eye closure, V2 and V3 have strongly attenuated 421 BOLD regardless of temporal contrast. Red plots indicate high temporal contrast trials and blue 422 indicates the low temporal contrast trials. The grey bar indicates 16 second period of stimulus 423 presentation. Error shading is 1 SEM. N=8 subjects. (B) Average BOLD activation during stimulus 424 presentation across conditions. Pairwise comparisons show a significant decrease in V2 and V3 BOLD 425 magnitude with eye closure for high temporal contrast stimuli. In LGN, BOLD magnitude with high 426 temporal contrast stimuli with eyes open was marginally greater than low contrast (t(14)=1.79; P = 427 0.047) at a Bonferroni corrected p-value cutoff of 0.0125. In V1 with eyes closed, BOLD magnitude 428 During high temporal contrast stimuli was marginally greater than during low temporal contrast stimuli 429 (t(14)=1.70; P=0.055). In V2, BOLD magnitude with high temporal contrast stimuli with eyes open 430 was greater than low contrast (t(14)=2.51; P=0.012) and high temporal contrast stimuli with eyes 431 closed was suppressed compared to eyes open. In V3, BOLD magnitude during high temporal contrast 432 stimuli with eyes closed was also suppressed compared to eyes open. Y-axis is BOLD signal averaged 433 across 4-16s post-stimulus onset. Error bars are 1 SEM. All p-values from pairwise comparison only 434 survive multiple comparison correction at a p-value less than 0.0125, using Bonferroni correction 435 (0.05/n where n=4 per ROI). * *P* < 0.0125 436

- 437 Figure 3. The effects of eye closure do not depend on eccentricity tuning. (A) Event-triggered
- 438 Average for BOLD response to luminance task across cortical ROI and eye condition separated by
- 439 voxels tuned to different portions of the visual field. With eyes open and eyes closed, the BOLD
- 440 responses to high contrast stimuli are uniform across eccentricities for all cortical ROIs. Foveal voxels
- 441 were tuned to between 0.01 dva 1.5 dva. Parafoveal voxels were tuned to between 1.5 dva 4.0
- 442 dva. Peripheral voxels were tuned to between 4.0 dva 17 dva. (B) Average BOLD activation during
- stimulus presentation across conditions (top = eyes closed; bottom = eyes open), separated by
- 444 eccentricity preference. There are no significant pairwise comparisons when comparing eccentricity
- responses within each ROI. Y-axis is BOLD signal averaged across 4-16s post-stimulus onset. Error
- bars and error shading is 1 SEM.

Eye closure produces distinct responses in early and late visual processing pathways

Methods

Subjects viewed a temporal contrast modulated visual stimulus with eyes open or eyes closed while collecting fMRI



Outcomes



Conclusions

Visual information is processed in early visual processing regions with eyes open and closed, whereas eye closure attenuates visual responses in later visuocortical regions

Sample stimulus blocks







