Vestibular Activation Differentially Modulates Human Early Visual Cortex and V5/MT Excitability and Response Entropy

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Head movement imposes the additional burdens on the visual system of maintaining visual acuity and determining the origin of retinal image motion (i.e., self-motion vs. object-motion). Although maintaining visual acuity during self-motion is effected by minimizing retinal slip via the brainstem vestibular-ocular reflex, higher order visuovestibular mechanisms also contribute. Disambiguating self-motion versus object-motion also invokes higher order mechanisms, and a cortical visuovestibular reciprocal antagonism is propounded. Hence, one prediction is of a vestibular modulation of cortical excitability and indirect measures have variously suggested none, focal or global effects of activation or suppression in human visual cortex. Using transcranial magnetic stimulation-induced phosphene thresholds to probe cortical excitability, we observed decreased V5/MT excitability versus increased early visual cortex (EVC) excitability, during vestibular activation. In order to exclude nonspecific effects (e.g., arousal) on cortical excitability, response specificity was assessed using information theory, specifically response entropy. Vestibular activation significantly modulated phosphene response entropy for V5/MT but not EVC, implying a specific vestibular effect on V5/MT responses. This is the first demonstration that vestibular activation modulates human visual cortex excitability. Furthermore, using information theory, not previously used in phosphene response analysis, we could distinguish between a specific vestibular modulation of V5/MT excitability from a nonspecific effect at EVC.

Keywords: human V5/MT, response entropy, TMS, vestibular

Introduction

Much of what we know of the cerebral cortical function underlying human visual perception has been obtained during the head static condition. Ecologically however, the visual system is more accustomed to functioning during head motion, and here a visuovestibular interaction is required to optimize visual acuity and to resolve self-motion versus object-motion ambiguity. The problem for the visual system is exemplified by the situation of climbing a tree with the aim of picking ripe fruit. Successful fruit picking requires adequate visualization of the fruit despite motion of oneself or the fruit. The task would not be considered a success if we fell out of the tree, and to avoid this, it is imperative that we distinguish between our own motion (passive or active) from that of the swaying branches (from which we might infer, sometimes erroneously, our own movement). The vestibular system contributes to the visual processes used in locating the fruit and avoiding falling by 1) stabilizing gaze and 2) helping to resolve self-motion from background motion (of the branches). Conventionally, visuovestibular interaction is thought of primarily as a brainstem process; first during self-motion, the vestibular-ocular reflex (VOR) improves visual acuity by stabilizing gaze upon the fovea, and second, visual inputs synapse upon primary vestibular neurons (Waespe and Henn 1977) to enable the detection of low frequency or constant velocity self-motion since the vestibular system is only sensitive to head accelerations.

Despite the eloquence of brainstem visuovestibular interaction, sensory signals may conspire to defeat a correct brain decision regarding self-versus object-motion and thus engender false sensations of self-motion (called 'vection'); for example, we may erroneously perceive ourselves to be swaying when faced with the swaying of branches when climbing a tree. One solution for resolving the self-versus object-motion problem is to invoke a reciprocal antagonism between visual and vestibular signals of motion. Evidence for a visuovestibular reciprocal inhibition is derived from psychophysical experiments assessing perception of self- or object-motion during concurrent visuovestibular stimulation (Probst et al. 1985, 1986) and at cerebral cortical level by reduced visual cortex responses during vestibular stimulation in functional imaging (Wenzel et al. 1996; Bense et al. 2001; Bottini et al. 2001; Stephan et al. 2005; Dieterich and Brandt 2008) and visual evoked potentials (Probst and Wist 1990). Despite conflicting evidence on visual cortical activation (some imaging and neurophysiological studies showing no effect of vestibular stimulation; Iida et al. 1997; Lobel et al. 1998; Suzuki et al. 2001; Engelhardt et al. 2007), an inhibition of visual motion cortex function could be functionally beneficial during excessive vestibular stimulation, by attenuating disorientating visual/self-motion percepts (Seemungal et al. 2011). Conversely, suppressing early visual cortex (EVC), which includes V1, could have wider, potentially inimical effects on visual processing since EVC is the primary entry route for visual signals to the cerebral cortex. Additionally EVC may be sine qua non for the conscious awareness of all visual percepts, including motion (Pascual-Leone and Walsh 2001; Silvanto et al. 2005). Indeed there is evidence for a higher order contribution to the maintenance of visual acuity above that explained by VOR gaze stabilization, during vestibular activation (Guedry and Ambler 1973; Guedry 1974; Tong et al. 2006). Thus a simple visual cortical inhibition could compromise visual discrimination during self-motion. To resolve the apparently competing brain excitation-suppression requirements between visual discrimination and motion processing during vestibular activation, one solution for the brain would be a selective down-modulation of visual motion areas, specifically V5/MT (Zeki 1974; Born and Bradley 2005), during vestibular activation. Current data showing a vestibular-mediated...
suppression of visual cortex are disparate however, suggesting variously a global suppression of visual cortex including V1 (Wenzel et al. 1996; Bottini et al. 2001) or focal suppression at V5/MT (Bense et al. 2001) or even V5/MT activation (Fasold et al. 2002).

Critically direct measures of human visual cortical excitability during vestibular activation have not been previously reported. Here, we describe the use of single pulse transcranial magnetic stimulation (TMS) to probe visual cortical excitability during vestibular activation (see Materials and Methods and Fig. 1). Change in cortical excitability in response to a sensory input is not, however, sufficient to establish specific functional relevance between stimulus and response, for example, cat primary visual cortex neuronal excitability was equally potentiated by vestibular and nociceptive inputs implying a nonspecific effect (Gorgiladze and Smirnov 1967). To further interrogate the functional relevance of excitability changes accompanying vestibular activation, we assessed changes in 1) phosphene percept—specifically phosphene size and intensity and 2) response entropy derived from an information theoretic analysis (see also Supplementary Materials). We first assessed TMS responses in area V5/MT since this region is critically important in visual motion processing (Zeki 1974; Born and Bradley 2005). We then assessed EVC responses during vestibular activation since, if vestibular signals modulate V5/MT excitability, one question is whether such modulation occurs via V1-dependent or V1-independent pathways. Certainly, modulation of human EVC excitability by magnetic or electrical stimulation (Pascual-Leone and Walsh 2001; Antal et al. 2003) induces parallel changes in V5 excitability. Recent reports (Schmid et al. 2010), however, have confirmed the existence of direct subcortical inputs to V5/MT bypassing V1, thus excitability changes in V5/MT divergent from those in EVC could allude to the pathways by which vestibular signals could modulate visual cortex.

Materials and Methods

Subjects

All 20 subjects gave informed consent prior to participating in the study, which had been approved by the local National Health Service Research Ethics Committee (there were 12 subjects per experiment with 4 subjects participating in both experiments). All subjects had no prior medical history (including ear problems or vertigo) nor were they on regular medication. Safety precautions including relevant subject exclusion criteria (e.g. excluding subjects with a personal or family history of epilepsy) utilized published recommendations (Rossi et al. 2009).

Following consent but prior to inclusion, subjects were tested to ascertain whether they could reliably perceive phosphenes in both hemispheres (see below for the technique used). Next, subjects were required to tolerate caloric irrigation with no ill effects (e.g., nausea). Six subjects in Experiment 1 and 5 subjects in Experiment 2 had already experienced caloric irrigation with documented normal nystagmic and vertigo responses. Caloric-naïve subjects were given 2 cold water irrigations on a day separate to the actual experiment to ensure both normal vestibular functioning and tolerability for the subject.

Using TMS to Probe Visual Cortical Excitability

Electrical field stimulation of the visual cortex, including that by TMS, produces illusory percepts of light called "phosphenes" (Brindley and Lewin 1968). The probability of evoking a phosphene is related to the instantaneous visual cortical excitability (Aurora and Welch 1998; Boroojerdi et al. 2000; Rauschecker et al. 2004; Romci et al. 2010). Thus the TMS intensity required to elicit a phosphene can be used to probe visual cortical excitability at the location of the applied magnetic pulse. Our experimental strategy consisted first of determining an individual’s 50% phosphene threshold (i.e., the TMS intensity which evoked a phosphene 50% of the time) at a given cortical location using a modified binary search algorithm—"MOBs" (Tyrell and Owens 1988). We chose a 50% phosphene detection rate since 1) many previous papers have used a 50% phosphene detection rate as a measure of visual cortical excitability (e.g., Aurora and Welch 1998; Boroojerdi et al. 2000), so our use of a similar measure allows an easier comparison with previous data; 2) a 50% detection rate allowed us to observe equally, increases or decreases on rate of phosphene detection with our intervention (i.e., vertigo).

Thus, using the same TMS intensity as that used to obtain a baseline 50% phosphene detection rate, we recorded the probability of inducing phosphenes during vestibular activation via caloric irrigation and twice more during the postcaloric period. This sequence was carried out 4 times in a subject, that is, twice for each ear and each hemisphere (Fig. 1). We assessed the change in cortical excitability with vestibular activation in 12 healthy volunteers (average age 28 years, 6 males) in area V5/MT (Experiment 1) and then in EVC (Experiment 2), in 12
TMS Stimulation Parameters
TMS was delivered to the cranial region of interest using a Magstim 200 stimulator (Magstim Co., UK) via a 70 mm figure-of-eight-shaped coil held in place by a dedicated coil holder. V5/MT stimulation sites were located using a functional method typically used in studies investigating phosphene perception (Aurora and Welch 1998; Guzman-Lopez et al. 2011; Schwarzkopf et al. 2011; and for a detailed discussion, see Walsh and Pascual-Leone 2003). A point was marked on the cranium, 3 cm dorsal and 5 cm lateral to the inion. Stimulation (at 80% of stimulator output) commenced at this point, and the coil was then moved around this spot (up to a maximum of 0.5 cm from the marked point) until a location from which a consistently vivid phosphene was induced. Eleven subjects saw clear and consistently moving phosphenes (one subject reported some variability in seeing the phosphenes move) in both hemispheres at this location, either at phosphene threshold or at a higher TMS intensity. For V5/MT, average phosphene thresholds, expressed as a percentage of maximal stimulator output, were as follows: all thresholds—63.1%; right and left hemispheres—60.6% and 65.5%, respectively; ipsilateral and contralateral conditions—63.3% and 62.8%. EVC was determined by measuring 2 cm dorsal and 1 cm lateral to the inion. The coil was moved around an area of maximum radius 0.5 cm from the marked point until a vivid phosphene was obtained. No moving phosphenes were seen at this location. For EVC, average phosphene thresholds were as follows: all thresholds—68.6%; right and left hemispheres—68.6% and 68.5%, respectively; ipsilateral and contralateral conditions—68.1% and 69.0%.

Caloric Irrigation
Subjects underwent cold water caloric irrigation in the prone position to near 30 °C below the horizontal (Fig. 1). Following otoscopy, the external auditory meatus was irrigated with water at 30 °C at a rate of 500 mLs/min for 40 s. The onset of vertigo and slow-phase vestibular nystagmus typically reaches a maximum 20 s after the end of the irrigation and lasts up to 2 min (Hood and Korres 1979). Subjects’ nystagmic responses were not measured during the phosphene experiments, but we confirmed that subjects experienced dizziness as in preexperiment caloric irrigations.

Entropy Estimation
Entropy is a measure of the uncertainty associated with a random variable. In the case of an equally weighted coin being flipped, the uncertainty associated with the outcome would be 1 bit—the maximum entropy possible for 2 outcomes. It was shown by Shannon (1948) that entropy is the maximum correct way to measure this uncertainty; it is therefore of interest to examine phosphene probabilities in entropy terms, as this allows us to directly interpret measured changes in phosphene probability as a reduction in the uncertainty of the phosphene report. To assess how much the uncertainty of phosphene reporting was reduced by caloric irrigation, we measured the yes/no response of subjects to each of 80 TMS pulses per experimental run. There were 4 runs per subject, thus this sequence of 80 pulses was repeated 4 times for each subject. In order to quantify the entropy over trials, we first calculated the probability of a yes response for each run (over subjects and repeats). This probability distribution \( p \) was used to calculate the Shannon entropy (Shannon 1948):

\[
H = -\sum_{i=1}^{K} p_i \log_2 p_i \tag{1}
\]

where the index \( i \) runs over \( K \) possible trial outcomes. In our case, \( i \) can be 0 (no phosphene) or 1 (phosphene observed). In practice, we estimated \( H \) using the NSB entropy estimation algorithm (Nemenman et al. 2004), a Bayesian technique using a Dirichlet prior, as described in Supplementary Materials. All entropy calculations were bootstrapped 200 times and have been presented as mean ± standard deviation over this bootstrapped distribution (i.e., standard error of the entropy) in the text.

Results
Effect of Vestibular Activation on Probability of Perceiving Phosphenes
Vestibular activation significantly reduced the probability of perceiving V5/MT phosphenes as compared with baseline (\( P < 10^{-4} \), Binomial test; Fig. 2A). In contrast, vestibular activation “increased” the probability of perceiving V1 phosphenes, although the magnitude of the effect was less than that seen in V5/MT (\( P = 0.002 \), Binomial test; Fig. 2A). We also analyzed the phosphene responses via a repeated measures analysis of variance (ANOVA) with factors “cortex” (2 levels: V5/MT vs. EVC) and phase (4 levels: viz, “baseline,” “vertigo,” “recovery1,” and “recovery2”). Given the pattern of response shown in Figure 2A, the lack of significant (\( P > 0.05 \)) main effects of cortex and “phase” were to be expected, but the significant interaction between cortex and phase (\( F_{3,9} = 4.24, P = 0.04 \)) confirmed the dichotomous response pattern between V5/MT versus EVC. An analysis of the individual responses (Fig. 2B) showed that for V5/MT, 8 subjects perceived fewer phosphenes during vestibular activation and only 2 subjects showed a prominent increase in phosphene frequency. The pattern for EVC differed from that for V5/MT with most subjects showing increases in perceiving EVC phosphenes during vestibular activation.

To illustrate the time course of the effect of vestibular activation on visual cortical excitability, the binary responses (“yes” and “no”) were converted into a “probability score” with yes (1) and no (0) responses summed to produce a score for every 5 stimuli. Thus the 20 responses per phase were binned into 4 epochs. Figure 2C shows the average scores obtained for each of the 4 epochs per phase across all subjects. The binned data displays the clear divergence between the scores for V5/MT (attenuation) compared with V1 (augmentation) underlying changes in cortical excitability in response to vestibular activation.

When the effect of vestibular activation from Experiment 1 (V5/MT phosphenes) was divided into ipsilateral (irrigation on same side as TMS stimulation) and contralateral, the effect of vestibular activation on attenuating phosphene perception was still significant (Fig. 2D; Bonferroni corrected significance level \( P=0.01 \)), although the size of the effect was more prominent in the ipsilateral condition. Similarly, we found a significant effect of vestibular activation on attenuating the probability of perceiving right and left hemisphere elicited V5/MT phosphenes (Fig. 2D) with a more marked effect in the right hemisphere. The same analysis for EVC found no significant difference in the size of the effect between hemispheres.

Effect of Vestibular Activation on Phosphenes Percept
Subjects were told to give a rating to the phosphenes obtained during baseline of 3/5 for SIZE and INTENSITY and then to give subsequent phosphene ratings relative to baseline (and of 5) at the end of each phase for each run. The averaged data for SIZE and INTENSITY responses are shown in Fig. 2E demonstrating that vestibular activation modulated V5/MT but not V1 phosphene characteristics. We analyzed phosphene SIZE and INTENSITY ratings in response to vestibular activation for both EVC and V5/MT, in 2 ways:
1. First, we assumed that subjects maintained an accurate memory of the baseline rating (i.e., 3/5). We thus performed a 1-sample t-test between rating scores obtained for the “vestibular activation” condition as compared with a hypothetical mean of 3 (of 5). This showed significant effects for V5/MT for SIZE ($t = 2.73, df = 47, P < 0.01$) and INTENSITY ($t = 3.64, df = 47, P < 0.001$) but not for EVC ($P > 0.05$) for either rating (SIZE or INTENSITY).

2. The second method assumed that the recovery back toward baseline from the vestibular activation condition more...
accurately described subjects' internal rating of phosphene SIZE and INTENSITY. Here, we performed separate repeated measures ANOVA for INTENSITY and SIZE estimates with factors “laterality” (ipsilateral versus contralateral) and “phase” (vestibular activation, Recovery 1, Recovery 2). For V5/MT, phosphene INTENSITY showed a significant main effect of laterality ($F_{1,23} = 9.57, P = 0.005$) and phase ($F_{2,22} = 3.53, P < 0.05$) but with no interaction between laterality and phase. For EVC, we found no significant effect of vestibular activation on perceived phosphene SIZE or INTENSITY. In summary, the data show that vestibular activation attenuates V5/MT phosphene INTENSITY; the evidence for an effect on phosphene SIZE at V5/MT was however less compelling. In contrast, all the analyses showed no effect of vestibular activation on EVC phosphene characteristics.

Information Theoretic Analysis of Effect of Vestibular Activation on Visual Cortical Excitability

We found that vestibular activation affected the excitability of both EVC and V5/MT, but this does not necessarily prove that vestibular signals have a specific functional interaction with visual cortex. To further interrogate the specificity of the vestibular effect on phosphene reports, we hypothesized that a functionally relevant visuovestibular interaction would be indicated by a significant reduction in the uncertainty associated with phosphene reports as assessed by an entropy analysis.

We titrated the baseline TMS intensity to yield a phosphene response rate of close to 50%, that is, the probability of perceiving a phosphene was 0.5, which, for a binary choice situation, is the state of maximum uncertainty (or maximum “Entropy” = 1 bit). We obtained the response entropy (pooled across subjects and repeats) throughout the 80 TMS stimulus sequence (Fig. 3) for baseline (stimuli 1–20), vestibular activation (stimuli 22–40), and recovery phases (stimuli 41–60 and 61–80). As the simple “plugin” estimate of entropy is biased by limited sampling (Miller 1955; Panzeri and Treves 1996), we used the NSB entropy estimator (Nemenman et al. 2004) for our calculations. Using surrogate data, we were able to demonstrate that the expected bias using this estimator with our sample set size (80 trials) is limited to a few percent (see Materials and Methods and Supplementary Materials). The key

![Figure 3](https://academic.oup.com/cercor/article-abstract/23/1/12/327261)

**Figure 3.** Response entropy. The 4 panels (V5/MT on the left and EVC on the right) relate to response data pooled from the subjects for each TMS stimulus in a run and ordered in the sequence of presentation; that is, each run consisted of 4 phases and 20 stimuli/phase so there were 80 TMS pulses per run (x-axis of panels = 1–80). The top panels show the pooled $P_\text{A}$ (probability of seeing a phosphene), and the bottom panels show information entropy ($H$). For V5/MT, the baseline entropy is stable but a clear decline in $H$ occurs with vestibular activation. In contrast, for EVC, the moderate decline in $H$ with vestibular activation is comparable to the entropy spikes that seem to occur spontaneously throughout the record.
observable effect was a significant reduction in entropy for V5/MT stimulation responses during the 21st and 22nd trials (from 0.96 ± 0.03 bits at baseline to 0.73 ± 0.10 bits in the 22nd trial, significant at \( p \approx 10^{-140} \) with two-sample Kolmogorov-Smirnov test), corresponding to the onset of vestibular activation.

**Discussion**

Our main finding was that vestibular activation resulted in opposing effects on phosphenes reports at area V5/MT versus EVC, specifically, a reduction at V5/MT versus an increase in EVC phosphenes reports were observed during vestibular activation.

**Specific and Nonspecific Effects of Vestibular Activation on Visual Cortex**

Our results additionally suggest a specific vestibular effect on visual cortex area V5/MT. First, that EVC and V5/MT results were divergent excludes a generalized effect of vestibular activation (e.g. arousal) on visual cortical excitability. Second, using a similar protocol, we have recently shown that visual motion adaptation increases V5/MT excitability (Guzman-Lopez et al. 2011). Third, V5/MT phosphenes intensity and size were attenuated by vestibular modulation. Note that the task required subjects to both detect phosphene and assess their characteristics (i.e., phosphenes size and intensity) and thus divided their attention. This is a potential reason for attenuating phosphenes percepts, but this was the case for all phases, including baseline (where subjects were also required to focus on phosphenes presence and characteristics in applying a 3/5 size and intensity score) and was the same for both V5/MT and EVC phosphene where we obtained opposing responses. Fourth, that we found a larger vestibular effect on cortical excitability in the right hemisphere is consistent with reports of a right hemisphere bias for vestibular cortical processing (Dieterich et al. 2003; Seemungal et al. 2008) and provides further evidence against a nonspecific effect for our results. Last, we found that V5/MT response entropy (Fig. 3) changed markedly during the initial period of vestibular activation (where vestibular activation is maximal), in stark contrast to highly stable response entropy in the pre- and postvestibular epochs.

Theoretically, caloric-evoked eye movements could contribute to the observed vestibular effects on V5/MT excitability (i.e., suppressive), since vestibular effects were more prominent when the fast phase was directed toward the stimulated (with TMS) hemisphere. Such an eye movement-related effect could not however account for "all" of our findings since we also found significant inhibitory effects when the evoked nystagmus was directed away from the stimulated hemisphere. Additionally, 2 previous studies (Thilo et al. 2004; Boulay and Paus 2005) failed to find an eye-movement modulation of visual cortical excitability (measured by TMS-induced phosphene).

Another issue is whether vertigo, which itself creates illusions of self-motion, could be more distracting to the perception of moving V5 phosphene than to static phosphene and thus explain the discrepancy in response between the V5 and EVC. An additional control experiment (see Supplementary Materials) where we assessed V5/MT phosphene reports during illusory self-motion engendered by a rotating sound source (called "auditory vection") failed to show any suppressive effect on phosphene (there was a trend for increased phosphene reports during vection). These data thus suggest that it is vestibular activation and not a sensation of self-motion per se that results in an attenuation of V5/MT phosphene reports.

Our data suggest that vestibular activation augments EVC excitability in a nonspecific manner. We found no effect of vestibular activation on reported phosphene size and intensity, and no laterality effect (right versus left hemisphere) on reported phosphene probability, implying a more general effect for EVC. EVC response entropy did dip slightly with vestibular activation but this appeared unremarkable in the context of repeated entropy "spikes" (Fig. 3) perhaps alluding to the multimodal sensory inputs to EVC. For example, vestibular, pain, and auditory stimulation all enhance V1 neuronal excitability (Gorgiladze and Smirnov 1967; Romei et al. 2007).

Single cell data in primates (Chowdhury et al. 2009) demonstrate a modulation of V5/MT neuronal activity by rotations in the light but with marked attenuation during dark rotations with a light point fixation. These data suggest an optic flow modulation rather than a direct vestibular drive to the sampled V5/MT neurons but do not however exclude the possibility that V5/MT neurons could show a vestibular modulation only during concurrent visual motion stimulation. Such a modulation was found in the lateral geniculate nucleus (LGN) (Papanioannou 1973), where the caloric effects on LGN light-evoked neuronal responses were independent of caloric effects upon spontaneous activity.

Although area V1 is the major input to V5/MT, hence explaining the finding that changes in human V1 excitability (e.g., by magnetic or electrical stimulation) generally match changes in V5/MT excitability (Antal et al. 2003; Silvanto et al. 2005), our data showing a dichotomous effect on EVC versus V5/MT excitability imply a vestibular modulation of V5/MT activity via direct subcortical pathways to V5. Indeed, there is recent evidence supporting direct LGN and collicular inputs to V5/MT (Sincich et al. 2004; Nassi and Callaway 2006; Berman and Wurtz 2008; Lanyon et al. 2009; Lyon et al. 2010; Schmid et al. 2010). This data allied with evidence for vestibular reactivity in both subcortical loci (Papanioannou 1973; Bisti et al. 1974; Magnin et al. 1974), provides a potential anatomical explanation for our findings.

**A Role for Vestibular Modulation of Visual Signals?**

One role for a vestibular modulation of visual motion signals at cortical level would be optimizing visual form perception. The magnocellular system conveying visual motion information to V5/MT via a low latency pathway (Bullier 2001; Born and Bradley 2005) could be suited to a vestibular modulation, since vestibular signals possess low latencies themselves. Beckers and Zeckii (1995) found evidence of a fast direct non-V1-dependent pathway relaying visual motion signals to V5/MT in humans. Furthermore, recurrent (and rapid) V5/MT-V1 pathways modulate the earliest components of striate neuronal responses that affect motion processing, including direction and motion selectivity responses (Hupé et al. 1998; Galuske et al. 2002). Such recurrent pathways could also benefit visual form processing if head motion predictive signals were utilized to optimize vision during head turns when minor degrees of retinal image slip and thus defoveation could occur. Supporting this notion is the observation in humans that visual discrimination is enhanced during vestibular activation (Guedry and Ambler 1973; Guedry 1974; Tong et al. 2006) and in animals that inactivating V5/MT impaired both visual motion and global
visual form responses in V1 (Hupé et al. 1998). It is tempting to speculate that vestibular signals, with latencies in the order of 5 ms, could specifically modulate those fast V1-independent pathways which could be employed in a motion predictive mechanism to enable a rapid retuning of visual perception to improve visual form perception. Indeed Vannier-Mercier and Magnin (1982) considered such a role for vestibular signals following their demonstration of a vestibular modulation of V1 neuronal activity in cats.

A role for vestibular signals in optimizing visual form perception does not obviate a vestibular role in resolving the self- versus object-motion conflict. Indeed, neuronal centre-surround inhibition, a prominent feature of V5/MT neurons, is one suggested neuronal mechanism (Huang et al. 2008) mediating motion perception discrimination (for review, also see Born and Bradley 2005). The classical neuronal receptive fields, where increased neuronal firing rate occurs in response to visual targets within a specific retinotopic location, are attenuated by centre-surround inhibition mechanisms (prominent in V5/MT), which are engaged by large visual targets. Thus small high-contrast stimuli would engender overall neuronal activation and large visual targets overall neuronal inhibition. Centre-surround inhibition may play a role in visual-vestibular phenomenon, such as vection, whereby large field visual stimuli (e.g., moving clouds) elicit sensations of self-motion. Vection carries the risk of an erroneous interpretation of self-motion. Centre-surround mechanisms, by modulating the integration or segmentation of a moving visual scene, may help resolve this object/self-motion ambiguity (Huang et al. 2008). Hence, the centre-surround mechanism could mediate the reported visual-vestibular antagonism suggested by human imaging studies (for review, see Dieterich and Brandt 2008).

Bimodal Sensory Interaction, State Dependent Excitability, and Information Content

Recent neurophysiological data demonstrate that bimodal stimulation engenders cortical neuronal responses with more information content than with unimodal stimulation (Kayser et al. 2010). Additionally, when the result of such bimodal combination is a reduction in neuronal activity, the information content is further increased. Indeed, we have recently quantified the state-dependency of TMS-induced V5/MT responses (Guzman-Lopez et al. 2011) showing that adaptation to a random dot visual motion stimulus induces a facilitation of TMS responses. Conversely, we suggest that vestibular activation might reduce the level of neuronal excitability in V5/MT by suppressing visually evoked activity that is incompatible with the vestibular cue, resulting in an overall more sparse V5/MT activity. In contrast, a nonspecific vestibular effect on area EVC would be equivalent to adding noise thus making EVC more likely to be excited by TMS.

Conclusion

In summary, we found that vestibular activation specifically suppresses V5/MT excitability but engenders a nonspecific increase in Early Visual Cortical excitability. Such visuovestibular cortical interaction could underlie the perceptual mechanisms that attempt to resolve visual motion from self-motion percepts while simultaneously optimizing visual form perception under natural conditions of vestibular activation such as during locomotion.

Supplementary Material

Supplementary material can be found at: http://www.cercor.oxfordjournals.org/

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Notes

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