Dissociation of Eye and Head Components of Gaze Shifts by Stimulation of the Omnipause Neuron Region

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Gandhi NJ, Sparks DL. Dissociation of eye and head components of gaze shifts by stimulation of the omnipause neuron region. J Neurophysiol 98: 360–373, 2007. First published May 9, 2007; doi:10.1152/jn.00252.2007. Natural movements often include actions integrated across multiple effectors. Coordinated eye-head movements are driven by a command to shift the line of sight by a desired displacement vector. Yet because extraocular and neck motoneurons are separate entities, the gaze shift command must be separated into independent signals for eye and head movement control. We report that this separation occurs, at least partially, at or before the level of pontine omnipause neurons (OPNs). Stimulation of the OPNs prior to and during gaze shifts temporally decoupled the eye and head components by inhibiting gaze and eye saccades. In contrast, head movements were consistently initiated before gaze onset, and ongoing head movements continued along their trajectories, albeit with some characteristic modulations. After stimulation offset, a gaze shift composed of an eye saccade, and a reaccelerated head movement was produced to preserve gaze accuracy. We conclude that signals subject to OPN inhibition produce the eye-movement component of a coordinated eye-head gaze shift and are not the only signals involved in the generation of the head component of the gaze shift.

INTRODUCTION

We interact with our environment by generating actions coordinated across multiple effector systems. Integrated movements of the eyes, head, torso, and/or limbs are used to perform activities such as driving a car (Land 1992), hitting a ball (Land and McLeod 2000), or even generating defensive postures (Graziano et al. 1997). Studies probing the neural bases of integrated motor control have typically considered the coordination of oculomotor subsystems (Fukushima et al. 2002; Gardner and Lisberger 2001; Keller et al. 1996; Walton and Mays 2003) or of eye movements with skeletal effectors like the head (Crawford et al. 2003; Guitton et al. 2003; Pélieisson et al. 2003) and the hand (Cohen and Andersen 2002; Courjon et al. 2004; Crawford et al. 2004; Stuphorn et al. 2000).

Reorienting the line of sight is often accomplished by coordinated eye-head movements (Bizzi et al. 1971). In general, the saccadic eye and head components are temporally coupled (Freedman and Sparks 1997b; Phillips et al. 1995; Zangemeister and Stark 1982a); the saccadic eye component initiates the gaze shift, and the onset of the head movement lags behind, at least in part, because of its heavier inertial load. When the initial eye-in-head position is deviated in the direction of the gaze shift (Becker and Jürgens 1992; Freedman and Sparks 1997b; Fuller 1996) or when the location and timing of the stimulus are predictable (Bizzi et al. 1972; Moschner and Zangemeister 1993; Zangemeister and Stark 1982a,b), however, the lag in the head component can be reduced or it can even lead gaze onset. Furthermore, long-duration, subthreshold microstimulation of the superior colliculus, a structure believed to issue a desired gaze-displacement command (Freedman and Sparks 1997a; Freedman et al. 1996; Klier et al. 2001), can evoke a head-only movement—the eyes counter-rotate in the orbits—or a head-only movement that precedes the onset of a gaze shift (Corneil et al. 2002; Pélieisson et al. 2001). Thus the timing of head initiation relative to gaze onset need not be rigidly constrained.

Independent control of the eyes and head occurs naturally at the level of the extraocular and neck motoneurons, although some form of separate processing has also been proposed to occur in cortical (Chen 2006; Chen and Walton 2005; Constantin et al. 2004, 2006), subcortical (Gandhi and Walton 2006; Pathmanathan et al. 2005a,b; Walton and Gandhi 2006), and cerebellar (Quinet and Goffart 2005) structures. It has been hypothesized that head movements associated with gaze shifts are driven by at least two parallel pathways: one is gated by the pontine omnipause neurons (OPNs) and issues movement commands to the eyes and head and the other bypasses the OPN gate and does not encode a saccadic eye-movement command (Corneil et al. 2002; Galiana and Guitton 1992; Goossens and Van Opstal 1997; Guitton et al. 1990). The objective of the present study was to test the functional contribution of head pathway(s) external to the OPN inhibition. Microstimulation was delivered to the OPN region either prior to or at the onset of horizontal head-unrestrained gaze shifts directed to the location of a briefly flashed target. Given that the OPNs inhibit excitatory burst neurons (Curthoys et al. 1984) that project to the abducens motoneurons, it was expected that the microstimulation would prevent generation of the saccadic eye component of gaze shifts. The key additional predictions were initiation of a head movement would consistently precede the gaze shift when microstimulation was delivered before gaze onset and an ongoing head movement would continue along its trajectory when stimulation was triggered after gaze onset.

Preliminary versions of this data have been published previously (Gandhi and Sparks 2000, 2001a; Sparks and Gandhi 2003; Sparks et al. 2001).

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METHODS

General procedures

Two juvenile rhesus monkeys (Macaca mulatta) weighing 5–8 kg served as subjects. All experimental protocols were approved by the Institute Animal Care and Use Committee at the Baylor College of Medicine and complied with the guidelines of the Public Health Service policy on Humane Care and Use of Laboratory Animals. All surgical procedures were performed in a sterile environment and under isoflurane anesthesia. Using Vetbond tissue adhesive, a search coil was attached to the sclera of one eye of each monkey. Another coil, a head-restraint post, and a stainless steel chamber were secured to the skull. The cylinder was slanted laterally in the frontal plane at an angle of 26° so that a microelectrode penetration through its center would reach the midline in the interaural plane 1–mm below interaural zero.

Post recovery, the animal was trained to sit in a customized chair fitted with neck, waist, and chest plates that minimized (approximately ±10°) but did not prevent trunk and body movements. On each experimental day, the chair with the animal inside it was canted to the laboratory. A micro-miniature laser (Edmund Optics; Barrington, NJ) was attached to the animal’s head implant. In addition, a hydraulic microdrive was secured to the chamber and a parylene coated tungsten microelectrode (Micro Probe; Gaithersburg, MD) was inserted into the brain. The animal with its chair was then placed in a magnetic field frame (CNC Engineering; Seattle, WA). Electromagnetic induction technique was used to measure eye-in-space (gaze) and head-in-space positions from the coils around the eye and in the head implant, respectively. Eye-in-head position was computed as the difference of gaze and head positions. See Gandhi and Sparks (2001b) for more details.

Experimental setup and behavioral tasks

When seating in the magnetic field frame, the animal faced a flat visual display consisting of 41 rows and 49 columns of tri-state light-emitting diodes (LEDs) that could light up red, green, or yellow. It was placed 72 cm from the center of the field coil, which corresponds to inter-LED spacing of 2° on a tangential plane. Maximum target eccentricity, with respect to straight ahead, was ±48° horizontally and ±40° vertically. All target eccentricities reported in this manuscript are in tangential coordinates and can be transformed into rotational coordinates as described previously (Gandhi and Sparks 2001b; Huebner et al. 1995).

While the electrode was being lowered into the paramedian pontine reticular formation (PPRF) region, the animal performed two randomly interleaved behavioral tasks. At the onset of each trial, a red LED and the microminiature laser were illuminated. When turned on, the laser module emitted a red beam of light onto the visual display panel. The animal was granted 2,000 ms to both fixate the red target and align the laser beam (i.e., the head) inside an imaginary, computer-controlled window (radius: 4–6°) around the LED location and then maintain these requirements for 700–1,000 ms. At the end of this step, the eyes are approximately centered in the orbits, and gaze and head positions are nearly equivalent. The remainder of the steps was dependent on the ensuing behavioral task.

In the memory-guided gaze shift task, a gaze target (yellow LED) was flashed for 500–1,300 ms, while the head target (red LED) and miniature laser remained on. Another 500–900 ms after the offset of gaze target, the laser and red LED were extinguished, which cued the monkey to initiate a gaze shift. If gaze position reached within an imaginary, computer-controlled window (radius: 6–10°) around the flashed target location in ≤500 ms, the LED was re-illuminated, and a final fixation of 1,500–2,000 ms triggered delivery of a liquid reward. A relatively large window size was necessary to account for the variability of endpoints for movements made in the dark (Goffart et al. 2006; White et al. 1994) and for greater hypometria associated with large movements (Becker 1991).

In the gap gaze shift task, the LED and the laser were extinguished after alignment of the head and gaze with the red target. After a gap of 200–500 ms, during which neither fixation condition could be violated, a gaze (yellow) target was illuminated at an eccentricity of 40–60° and remained on until the end of the trial. The monkeys were allowed 800 ms to direct gaze into a computer-controlled window (radius: 6–10°) around the yellow target location. To allow completion of the head movement, a final fixation of 1,500–2,000 ms was required before a liquid reward was delivered. A window constraint was not placed on the head position signal during acquisition of the gaze target.

Stimulation trials were introduced once the electrode penetration entered the PPRF region of the brain stem and OPNs were identified based on their discharge characteristics (cf. Gandhi and Keller 1999). The initial objective was to optimize the microelectrode location within the OPN region. On a subset of the gap trials (25–50%), microstimulation (10–30 μA, 200–300 Hz, 100–300 ms, 0.25-ms duration cathode pulses) was applied to the OPN region 100–200 ms after the offset of the red LED and laser. The gaze target was illuminated after the offset of the stimulation, and the animals were rewarded for acquiring the gaze target. If the tip of the stimulation electrode was located on the fringe of the OPN area where stimulation produces a constant velocity gaze movement (Cohen and Komatsuzaki 1972), the electrode was moved to eliminate this effect. This positioning effort also served to minimize stimulation-induced activation effects of nearby pathways (e.g., reticulospinal, tectospinal, and interstitialspinal) on head-movement generation.

On satisfactory positioning of the electrode, trials with stimulation triggered during the gap period were reduced to ~5% of all trials, but other stimulation tasks were introduced. On a subset of the trials, microstimulation (10–30 μA, 200–300 Hz, 100–300 ms, 0.25-ms duration cathode pulses) was triggered on gaze onset, which actually was detected on-line as deviation of either gaze or head position outside of its respective computer-controlled window constraint. On another fraction of interleaved trials, the microstimulation was delivered 100–200 ms after the cue to initiate the gaze shift (target onset in the gap task and fixation point offset in the memory-guided gaze shift task). This trial type was designed to test whether and how the latencies of gaze and head movements are compromised by OPN stimulation prior to gaze onset. In this latter condition, if stimulation onset followed gaze onset, the trial was relocated to the “stimulation triggered on gaze onset” database. For both stimulation trials, the gaze target was extinguished with the onset of the stimulation. If gaze position reached within an imaginary, computer-controlled window (radius: 6–10°) around the gaze target location in ≤500 ms after stimulation offset, the LED was re-illuminated, and a final fixation of 1,500–2,000 ms triggered a liquid reward.

In general, stimulation within the context of gap trials was more common at the beginning of each experimental day, as the electrode position within the OPN region was being fine tuned. On satisfactory placement of the electrode, the memory-guided gaze shift task, with and without stimulation, constituted ~75–80% of all trials. The rest of the trials were composed of the gap task with and without stimulation. Collectively, stimulation was delivered in 25–40% of all trials. Stimulation parameters were kept constant within a block of trials, irrespective of whether the current was injected before or after gaze onset. We typically only modified the stimulation duration in between blocks, when we felt it was necessary to prolong the stimulation-induced interruption to better visualize the effect of stimulation on the head component of the gaze shift. For the analyses, however, we did not parse the data according to stimulation duration as we did not consistently have enough trials for a specific stimulation duration and the same target configuration (as described in the following text).

A qualitative assessment indicated that stimulation produced similar effects on both gap and memory-guided gaze shift trials and hence
the data were pooled. Control data for the two trials types were also combined, as our gap trials did not produce an appreciable number of express latency gaze shifts. This is most likely due to factors such as small percentage of gap trials, variable gap durations and numerous target locations.

All target configurations required horizontal gaze shifts. The vertical eccentricity of all targets was 0° and, based on visual inspection, at eye level. Typically, the initial red LED was presented at 0, +20° (right visual field) or −20° (left visual field), and the yellow, gaze target appeared at ±40 or ±60° with respect to the initial target. If the gaze target location was beyond the visual display boundary, the most distal target (±48°) was illuminated to prevent any bias the animal might develop to evoke centripetal gaze shifts when the initial head and gaze positions were directed away from the midline. Gaze shifts directed to targets illuminated at 40 and 48° in the same hemifield were combined. Trials in which both initial and gaze targets were presented in the same hemifield (e.g., from 20° right to 40° right) were excluded from analysis.

Data acquisition and analyses

Data were acquired at 500 Hz using in-house software. Data were analyzed off-line using in-house software and Matlab (The Mathworks, Natick, MA). Typically, velocity criteria were used to detect the onset and offset of gaze, eye and head movements. For gaze and eye movements, the onset and offset velocity thresholds were 80 and 60°/s, respectively. For head movements, the onset and offset velocity thresholds were 8 and 5°/s, respectively.

The majority of the analyses presented in the manuscript required computing static parameters (latency, amplitude, magnitude, and time of peak head velocity and acceleration) and comparing them between control and stimulation conditions. In general, the Wilcoxon rank sum test—equivalently, the Mann-Whitney U test—that compares the medians of the distributions of control and stimulation conditions was used to test for statistical significance. When appropriate (and as indicated in the results), a pair-wise comparison was performed with the Wilcoxon sign rank test of equality of medians across matched samples of control and stimulation trials. For comparison, analogous t-test were also performed and comparable results were obtained. Additionally, a two-way ANOVA test was performed to determine whether any changes observed in the static parameters listed above could be statistically related to biomechanical factors or trial condition (stimulation or control).

A dynamics analysis was used to test the null hypothesis that stimulation did not significantly attenuate the head movement. For these analyses, all acceleration and velocity traces were aligned on head onset. A one-tailed t-test, modified to account for unequal variances in the control and stimulation trials (Milton and Arnold 1995), was applied to compute a P value for each 2-ms increment. The goal was to determine whether and when (relative to head onset) the head velocity in the stimulation trials was significantly attenuated. A significant attenuation in the head velocity and acceleration for ≥25 ms consecutively was required to count as a valid stimulation effect. The criterion of P < 0.05 was used for statistical significance across all tests.

RESULTS

Microstimulation was delivered to 23 sites within the OPN region of two monkeys (n = 13, monkey CH; n = 10, monkey BE) performing head-unrestrained gaze shifts. The stimulation was either delivered prior to movement initiation or triggered on the onset of the gaze shift. Data were sorted into six target configurations: rightward 40 and 60° gaze shifts initiated from the left hemifield (−20°), leftward 40 and 60° shifts from the right hemifield (20°), leftward gaze shifts (pooled 40 and 48°), from straight-ahead (0°), and rightward gaze shifts (pooled 40 and 48°) from straight-ahead (0°). The six target configurations and two stimulation modes (before and after gaze onset) produced a maximum of 12 different datasets to be analyzed for each stimulation site. After imposing the constraint that ≥6 trials must exist in each condition, 115 datasets were available for analysis: 48 for stimulation before gaze onset and 67 for stimulation after gaze onset. The eyes were approximately centered in the orbits prior to the onset of gaze shifts for control (0.21 ± 1.92°, mean ± SD) and both stimulation (0.25 ± 1.82°) conditions. The distributions were not statistically significantly different (rank sum test, P > 0.05).

The format of the data presentation order is as follows: a qualitative assessment of the effects of stimulation of the OPN region based on temporal profiles of gaze, head, and eye-in-head signals; an examination of whether and how stimulation modified the latencies of the head and gaze components; an investigation of stimulation-induced alterations in the kinematics and dynamics, with particular emphasis on the head component; an indirect evaluation of the gain of the vestibuloocular reflex (VOR) during the stimulation-induced perturbation; and, finally, a brief, statistical consideration of effects of biomechanical factors in addition to that of the stimulation.

Effects of OPN stimulation

Figure 1 illustrates the effects of stimulation of the OPN region on generation of rightward, 60° gaze shifts; the displayed trials are a subset of one dataset. Horizontal amplitude (left) and velocity (right) of gaze, head, and eye-in-head components are plotted as a function of time for individual control (cyan) and stimulation trials (red). To visualize the effects of stimulation delivered prior to gaze onset (Fig. 1A1), the movements are aligned on the cue to initiate the gaze shift. For control trials, the average gaze latency was ~250 ms. For the five stimulation trials shown in the figure, stimulation onset was initiated 150 ms after the cue and lasted for 300 ms. The onset of the gaze shift was delayed until after the end of the stimulation train. A head movement, however, was initiated during the stimulation (Fig. 1A2) but at a longer latency compared with control trials as judged by a visual comparison of the velocity traces. Gaze remained stable during this period, implying that the eyes counter-rotated in the orbits by the amount of the head movement (Fig. 1A3) and that the VOR gain was near unity. After stimulation offset, a gaze shift composed of an eye saccade and a reaccelerated head movement occurred. Gaze accuracy was maintained as the difference in the total gaze amplitude between control and stimulation conditions was small (stimulation-control, −1.19 ± 1.96°; median: −1.08°) across the 48 datasets. In contrast, the total head movement was greater in the stimulation condition (stimulation-control, 1.56 ± 3.05°; median: 1.06°; P < 0.01, sign rank test), and the reacceleration likely contributed to the hypermetria.

Example movements for which stimulation was triggered on the onset of gaze shifts are shown in Fig. 1B. These traces, obtained from the same animal and stimulation site illustrated in Fig. 1A, are aligned on gaze onset. Microstimulation produced an interruption of the gaze shift (Coble et al. 1994; Paré and Guitton 1998; Sparks et al. 2002) that resembled an interrupted saccade observed when the head is restrained.
movement kinematics (data not shown). Appropriate data collected in the gap task was comparable to the modulation in peak velocity, at least of head-restrained saccades (Edelman and Goldberg 2000). We speculate that this effect is most robust during visually guided movements. The movements illustrated here were performed in the Sparks 2000. We do not show the dual-peak modulation reported previously (Freedman and Sparks 2000), and were collected from the same stimulation site in 1 animal. The constant gaze position during the interruption suggests that the VOR gain was near unity during the interruption, nearly perfectly compensating for the sustained head movement. After stimulation offset, a resumed gaze shift, composed of a coordinated eye-saccade and a reaccelerated head movement, was observed. Gaze accuracy was maintained because the difference in the total gaze amplitude between control and stimulation conditions was small (control-stimulation, $-0.61 \pm 2.78^\circ$; median: $-0.51^\circ$) across the 67 datasets. In contrast, the total head movement was greater in the stimulation condition (stimulation-control, $3.55 \pm 2.90^\circ$; median: $3.19^\circ$; $P < 0.001$, sign rank test), and the reacceleration likely contributed to the hypermetria.

**Latency distributions**

Figure 2 quantifies the changes in latencies of gaze, head, and head relative to gaze when stimulation was delivered prior to a movement. Gaze onset was significantly delayed compared with control trials both within and across all datasets (Fig. 2A; rank sum test, $P < 0.05$). The distributions of gaze latency across the 48 datasets between control [266 ± 30 (SD) ms] and stimulation (416 ± 89 ms) were statistically different (rank sum test, $P < 0.0001$). Like gaze reaction time, head latency was also delayed by the stimulation, and significantly so in 35/48 datasets (Fig. 2B; rank sum test, $P < 0.05$). The distributions of head latency across all datasets were statistically different for control (268 ± 43 ms) and stimulation (359 ± 104 ms) trials (rank sum test, $P < 0.0001$). Figure 2C shows the timing of eye-head coordination (head-gaze latencies) was significantly altered by the stimulation in 43/48 datasets. Across all 48 datasets, head onset followed gaze onset by 2 ± 27 ms in the control condition but preceded gaze by 57 ± 40 ms during stimulation trials (rank sum test, $P < 0.0001$).

When stimulation was triggered after gaze onset (data not shown), the onset times of gaze and head were comparable in the control and stimulation conditions. Across the 67 datasets, the distributions of gaze latency for control (263 ± 30 ms) and stimulation (257 ± 33 ms) conditions were not significantly different. Neither were the distributions of head latency (control: 274 ± 44 ms; stimulation: 268 ± 39 ms) or head-gaze latency (control: 11 ± 27 ms; stimulation: 11 ± 24 ms; $P > 0.2$ for all 3 rank sum tests).

**Displacement measurements**

The effect of OPN stimulation on the eye and head components of a gaze shift was first evaluated by performing a displacement analysis. Figure 3A illustrates the method used to compute the stimulation-induced displacements when stimulation was delivered prior to gaze onset. The three panels display...
schematics of gaze, head, and eye-in-head movements as a function of time from one dataset (i.e., they have the same target configuration). All traces are aligned on head onset (leftmost first vertical dashed line at time 0). Each panel shows an averaged control movement (thick cyan traces) and two individual stimulation trials (thin waveforms). The initial components of the stimulation traces are shown in dark blue, but they are switched to red and marked with vertical dashed lines at the onset of the gaze shifts that follow the stimulation. First consider the stimulation trial labeled ①. The duration from head onset to gaze onset is identified as the span between the leftmost and the middle vertical dashed lines, and the change is position during this interval is computed for both the averaged control movement (cyan) and the individual stimulation trial (blue and red). This measurement is performed on gaze, head and eye-in-head channels, as illustrated in the three panels in Fig. 3A. This same procedure was then repeated for every stimulation trial. The schematic of the trial labeled ② reveals that the duration from head onset to gaze onset varies between trials and hence so do the displacement measures for both control and stimulation conditions. Thus for each dataset, this procedure yields two distributions (control and stimulation conditions) for gaze displacement, two for head displacement, and two for eye-in-head displacement. Furthermore, the number of elements in each distribution equals the number of stimulation trials in the dataset. The procedure was repeated for each of the 48 datasets.

To generate the summary exhibited in Fig. 3B, first consider the control and stimulation distributions for the gaze channel. For each dataset, the mean gaze displacement from the control distribution was plotted as a function of the mean gaze displacement from the stimulation condition (top). Thus the summary shows data from 48 datasets for which stimulation was delivered before gaze onset. A sign rank test was also applied to each dataset to determine whether the medians of the matched control and stimulations distributions were significantly different. Datasets showing statistically significant difference are shown as circles and crosses otherwise. The average change in gaze position across all datasets for control (17.8 ± 10.6°) and stimulation (0.8 ± 0.7°) conditions was significantly different (rank sum test, P < 0.0001).

The same procedure was then applied to the control and stimulation distributions for head and eye-in-head displacements. The summary data are plotted in the middle and bottom panels of Fig. 3B. The eye-in-head displacement was significantly different between control and stimulation conditions in...
47 of 48 datasets. The average change in eye position across all datasets for control (12.5 ± 7.7°) and stimulation (−2.2 ± 1.9°) conditions was significantly different also (rank sum test, \( P < 0.0001 \)). In contrast, the change in head position between control and stimulation conditions was significant for only 21 datasets. The distributions of head displacements across all datasets for control (4.7 ± 4.1°) and stimulation (3.1 ± 1.9°) conditions tended toward a statistical significance difference (rank sum test, \( P = 0.054 \)). For the 21 datasets that were significantly different, the mean change in head position was 6.7 ± 4.5° for control movements and 3.7 ± 1.5° for stimulation trials.

A similar approach was taken for the displacement analysis when stimulation was triggered on the onset of gaze shifts. The three panels in Fig. 4A are schematized waveforms of gaze, head, and eye positions as a function of time for an average control movement (thick, cyan waveforms) and for two individual stimulation trials (thin traces) aligned on gaze onset (leftmost first vertical dashed line at time 0). The initial components of the stimulation traces are shown in dark blue, but they are switched to red and marked with vertical dashed lines at the onset of the resumed gaze shifts that follow the stimulation-induced interruption in gaze. The change in gaze position across this epoch was computed for each stimulation trial and the average control movement. Because this duration is different across stimulation trials (see trials labeled 1 and 2), the displacements for the control distribution are nonidentical values. For each dataset, this procedure produces two distributions that were compared with a sign rank test as described in the preceding text. The same analysis was also performed for head and eye displacements. For the eye component, however, if the mean control gaze shift was completed before the end of the interval over which the displacements were measured, then the maximum eye-in-head displacement spanning the gaze shift was used (Fig. 4A, diamond in bottom panel). Note that the analysis on the eye-in-head component does not compare the ocular saccade amplitudes but rather the change in eye position during the same intervals over which the head displacements were evaluated (except if the control movement had completed, in which case, we considered the saccade amplitude for the eye-in-head displacement measurement).

Figure 4B compares the mean changes in gaze, head, and eye positions in control and stimulation trials across all datasets. The change in gaze (top) was significantly different between control and stimulation trials in 66/67 datasets. The average change in gaze position across all datasets for control (36.2 ± 5.8°) and stimulation (19.7 ± 6.3°) conditions was significantly different (rank sum test, \( P < 0.0001 \)). Similarly, the eye position displacement (bottom) was significantly different between control and stimulation conditions in 66 datasets. The average change in eye position across all datasets for control (20.4 ± 5.6°) and stimulation (5.2 ± 6.8°) conditions was significantly different also (rank sum test, \( P < 0.0001 \)). In contrast, the change in head position between control and stimulation conditions was significant for only 29 datasets. Unlike the consistent stimulation-induced decrease in gaze and eye displacement, however, the head displacement could either increase (9/29) or decrease (20/29). The distributions of head displacements across all datasets for control (15.6 ± 5.7°) and stimulation (14.6 ± 5.4°) conditions were not statistically different (rank sum test, \( P = 0.26 \)).

Velocity and acceleration measurements

Although the displacement analysis showed a profound effect on gaze and eye displacements, it did not yield as convincing a result for the head component when stimulation was triggered prior to gaze shifts (Fig. 3). When stimulation was triggered on gaze onset, the displacement analysis failed to find any significant effect on the head component (Fig. 4). It is possible that an effect might have been obscured because of the larger inertia of the head relative to the eyeball (Zangemeister et al. 1981) and because of an ongoing and perhaps attenuated neuromuscular drive to the head during the OPN stimulation. Hence, we further considered the effect of stimulation on the head movement by analyzing velocity and acceleration profiles for control and stimulation trials. Gaze and eye-in-head waveforms were not considered for this analysis.
Figure 5 plots head velocity, aligned on its onset, as a function of time for individual control (cyan) and stimulation trials (blue/red). Each panel represents trials from one dataset. If the stimulation was delivered prior to gaze onset (Fig. 5, A and B), the portion of the head movement that occurred before gaze onset is shown in blue; the remainder of the trial is plotted in red. Note that the duration of the blue trace is equivalent to the interval spanned by the vertical dashed lines in Fig. 3A. When stimulation was triggered on the onset of gaze shifts (Fig. 5, C and D), the head-movement component that spans from initial gaze onset to resumption of gaze shift following the interruption is shown in blue, and the rest in red. This duration is equivalent to the period spanned by the vertical dashed lines in Fig. 4A. Figure 6 shows the acceleration profiles of the same trials and in same format as Fig. 5.

The data illustrated in the top row in Figs. 5 and 6 were chosen to provide examples of significant attenuation on the head movement for each stimulation condition, whereas the traces shown in the bottom panel demonstrate cases of no obvious modification from the stimulation. In fact, on some stimulation trials, the head velocity and acceleration was higher compared with control trials. This effect was not consistent across all trials within a given dataset, suggesting that the effect was not specific to a stimulation site or to a target configuration. The effect of stimulation on head movements was quantified by examining the dynamics and kinematics of both velocity and acceleration. For each 2-ms increment throughout the duration of the blue traces in Fig. 5 (50 ms before head onset to gaze onset), the instantaneous velocity for control and stimulation movements were compared with a one-tailed *t*-test that was modified to account for potentially unequal variances in the two distributions (Milton and Arnold 1995) (see METHODS). Stimulation delivered prior to gaze onset (48 datasets) significantly lowered head velocity and acceleration in 37 and 32 datasets, respectively. The mean ± SD (median) attenuation in head velocity and acceleration occurred 34 ± 35 ms (18 ms) and 32 ± 50 ms (19 ms), respectively, after the onset of the head movement. Stimulation triggered on gaze onset (67 datasets) significantly lowered head velocity and acceleration in 35 and 23 datasets, respectively. The mean ± SD (median) attenuation in head velocity and acceleration occurred 77 ± 54 ms (78 ms) and 75 ± 38 ms (74 ms), respectively, after the onset of the head movement. In general, attenuation in head movements occurred later and was less frequent when stimulation was triggered after gaze onset.

The kinematics analyses on the magnitude and time of peak head velocity was performed in the same way as the displacement analysis. For each stimulation trial, the magnitude and time of peak head velocity was computed over the trace spanned from head onset to gaze onset (dark blue traces in Fig. 5). These values were paired with the measurements obtained over the same interval of the average control movement, and a sign-rank test was applied to compare the medians across paired samples of control and stimulation distributions for each dataset. The identical procedure was then performed on head acceleration traces.

The magnitude of peak head velocity was significantly different between control and stimulation conditions for 34/48 datasets when stimulation delayed gaze onset (Fig. 7A). Across all datasets, the distributions of peak head velocity for control (83.4 ± 46.7 °/s) and stimulation (54.1 ± 30.6 °/s) were significantly different (rank sum test, *P* < 0.001). The magnitude of peak acceleration was significantly different for 23/48 datasets (Fig. 7B). And, like peak head velocity, the distributions of peak head acceleration for control (1,702.3 ± 1021.6 °/s²) and stimulation (1,172.1 ± 713.0 °/s²) were also significantly different (rank sum test, *P* = 0.002). When stimulation was triggered on gaze onset, 30 of 67 datasets showed a significant modulation in the peak head velocity measure (Fig. 7C). In a few cases, the stimulation significantly increased peak head velocity. Across all datasets, however, the change was quite small, <4% attenuation, and the distributions in control (121.0 ± 58.4°/s) and stimulation (110.1 ± 52.7°/s) conditions were not significantly different (rank sum test, *P* > 0.2). For peak head acceleration (Fig. 7D), only 15 datasets showed a significant change. Like peak velocity, the stimulation barely reduced peak acceleration, by <2%, and there was no statistical difference between the control (2,173.7 ± 1,898.8°/s²) and stimulation (2,019.8 ± 1,816.2°/s²) distributions (rank sum test, *P* > 0.4).

Caution is required when comparing the times of peak velocity and acceleration between control and stimulation conditions. An examination of the temporal profiles in Figs. 5 and 6 reveals why this is the case. When stimulation delayed gaze onset...
onset (Figs. 5A and 6A), the head movement was accelerating when the gaze shift was initiated. In other words, the time of peak head velocity (and acceleration) is approximately equal to the duration of the head-only movement (each blue trace). Now note that the cyan, control waveforms across the same interval are also increasing, which implies that the peak times will be very similar for the matched pairs between control and stimulation trials. To circumvent this confound, we appended to each stimulation trial, the 100 ms of data after gaze onset, i.e., 100 ms after each blue trace ends, and computed the peak time measure. We also performed this operation on the datasets collected when stimulation was triggered on gaze onset, although this problem was not common for the long-duration stimulation-induced interruptions.

The time of peak head velocity was significantly different for 27/48 datasets when stimulation was delivered before gaze onset (Fig. 8A). Across the datasets, the distributions of control (132 ± 20 ms) and stimulation (159 ± 34 ms) conditions were significantly different (rank sum test, P < 0.0001). The stimulation-induced shift in time to peak velocity was 27 ± 33 (SD) ms across the datasets. There was a significant effect in the peak acceleration time for 26 datasets (Fig. 8B). Across all datasets, the distributions of control (80 ± 30 ms) and stimulation (112 ± 36 ms) conditions were significantly different (rank sum test, P < 0.0001). The shift in time was 33 ± 36 ms across the datasets.

Stimulation triggered on the onset of gaze shifts delayed the time of peak head velocity in 36 datasets (Fig. 8C). Across the 67 datasets, the distributions of control (140 ± 27 ms) and stimulation (189 ± 87 ms) conditions were significantly different (rank sum test, P < 0.001), and the stimulation-induced shift in time was 48 ± 77 ms. There was a similar, significant effect in the peak acceleration time for 25 datasets (Fig. 8D). Across all datasets, the distributions of control (85 ± 37 ms) and stimulation (124 ± 81 ms) conditions were significantly different (rank sum test, P < 0.01). Stimulation delayed the peak time by 39 ± 80 ms across the datasets.

Counter-rotation gain during the interruption

The examples and analyses of Figs. 1, 3A, and 4A demonstrate qualitatively that gaze position remains nearly constant during all stimulation-induced interruptions. Because the head continues to move during this period, the eyes must counter-rotate in the orbits by the amount of the head movement. We...
quantified this observation by plotting for each dataset the average change in eye position as a function of the mean head displacement over a period from head onset to gaze onset when stimulation was delivered prior to gaze onset (Fig. 9A). This interval is marked by the vertical dashed lines in Fig. 3A. When stimulation was triggered on gaze onset, the mean changes in eye and head positions during the interval between initial gaze shift offset and resumed gaze shift onset were compared (Fig. 9B). Note that this interval is not identical to the duration marked by the vertical lines in Fig. 4A. For both stimulation conditions, the points fall along the negative unity slope (dashed) line.

We quantified this observation with a counter-rotation gain parameter that was computed as the negative of ratio of the eye and head displacements plotted in Fig. 9. This approach of computing the counter-rotation gain is an indirect way to evaluate the VOR gain, which is typically computed as the negative of the ratio of eye and head velocities (Leigh and Zee 1999). A gain close to one would imply that the VOR is fully active. When stimulation was delivered before gaze onset, the mean \( \pm \) SD (median) gain was 0.70 \( \pm \) 0.27 (0.75). Similarly when stimulation was triggered after gaze onset, the mean \( \pm \) SD (median) gain was 0.85 \( \pm \) 0.21 (0.91). Next we subtracted the magnitudes of eye and head displacements during the counter-rotation phase for each dataset and used a sign rank test to determine whether the median difference across all datasets was significantly different from zero (\( P > 0.1 \), stimulation before gaze onset; \( P > 0.8 \), stimulation after gaze onset). Thus these results indicate the eye-in-head and head displacements during the counter-rotation were roughly equal in magnitude and opposite in direction, and imply that the VOR gain is close to unity during the interruption produced by stimulation of the OPN region.

Effects of biomechanical factors

Both the length of a muscle and its velocity of contraction constrain the force generated by a skeletal muscle. Because our experiments required different initial head positions (either roughly centered on the body or deviated \( \pm 20^\circ \) to the right or left) and different head dynamics (40 or 60\(^\circ\) gaze shifts), we sought to determine whether these parameters could account for the effects observed during OPN stimulation. Unfortunately, head-velocity profiles associated with the same initial head position and same desired gaze shift did not display enough variability to test for a combined effect of muscle length and velocity of contraction. Hence we tested for effects of these parameters separately.

To investigate for a potential effect of head position on the kinematics, we performed a two-way ANOVA with factors initial head position (centered or deviated) and trial type (control or stimulation). Table 1 lists the \( P \) values for the parameters reported in the preceding sections. Consistent with the previous study (Corneil et al. 2001), initial head position influenced most parameters associated with head movements during both stimulation conditions. Note, however, that the effects of the two factors were mostly independent. There were no statistically significant interaction effects when stimulation was delivered prior to gaze onset. When the stimulation was triggered on gaze onset, significant interactions were noted only on the times of peak head velocity and acceleration (Fig. 8) and on the amplitudes of gaze and eye displacements (Fig. 4B).

To test for an effect of head velocity, we sought to study gaze shifts of very different amplitudes generated from the same initial head position. Our reasoning was that head movements, and therefore head velocities, associated with these gaze shifts are different enough to be considered for statistical (ANOVA) testing. Thus we only considered trials with desired gaze shifts of 40 and 60\(^\circ\) and initial head position at 20\(^\circ\) contraversive to the direction of the movement. We performed a two-way ANOVA with factors desired gaze shift (40 and 60\(^\circ\)) and trial type (control or stimulation). Table 2 lists the \( P \) values for the parameters reported in the preceding sections. Consistent with the initial head position effect, the desired gaze amplitude—and by extension, head velocity—and the trial type significantly influenced head-movement parameters such as displacement and magnitude and time of peak velocity and acceleration, but there were no interaction effects on the head component.

Discussion

Converging lines of evidence indicate that at least two partially independent pathways are involved in the neural control of coordinated eye and head movements and that different modes of active eye-head coordination exist. A series of early influential studies (Bizzi et al. 1971, 1972) demonstrated that the relative timing of the eye and head components and the patterns of neck muscle activity observed when directing gaze to an unexpected stimulus (triggered mode) is quite different from that observed when animals look to anticipated targets (predictive mode). In general, the onset of the eye and head components are more tightly coupled in the triggered mode, but the head can often lead for large gaze shifts in the predictive mode (Bizzi et al. 1972; Moschner and Zangemeister 1993; Zangemeister and Stark 1982a). Behavioral paradigms designed to specify separate gaze and head-movement goals (Goossens and Van Opstal 1997) and tasks using auditory stimuli (Populin 2006; Tollin et al. 2005) have also demonstrated that the eye and head components of the coordinated movement can achieve a degree of independence.

FIG. 9. An indirect evaluation of the counter-rotation gain, which presumably reflects the vestibuloocular reflex (VOR) gain, was assessed by comparing the change in eye position as a function of the head displacement traversed during the stimulation-induced interruption in gaze. A: when stimulation was delivered before gaze onset, the changes in eye-in-head and head-in-space positions were measured over the interval from head onset to gaze onset (the region marked by the vertical dashed lines in Fig. 3A). B: when stimulation was triggered on gaze onset, the measurements were made across the interval starting at the end of the initial gaze shift and ending at the onset of the resumed gaze shift. Each point represents the average changes in eye and head positions for each dataset.
The results of our OPN stimulation experiments support the partially independent pathways hypothesis as they demonstrate a dissociation of the eye and head components of gaze shifts, even those produced in the triggered mode. We found that stimulation of the OPN region before gaze onset typically delays the saccadic eye movement until stimulation offset, on the order of several hundred milliseconds, without preventing the head movement (Fig. 2, A–C). Triggering the stimulation on the onset of the gaze shift arrests the ongoing saccadic eye movement without stopping the head movement. Furthermore, the duration of the head movement is quite different from the durations of the saccadic eye component and the gaze shift (data not shown, but qualitatively obvious). With early stimulation, for example, the head begins before gaze (and eye saccade) onset and continues after the gaze shift (and eye saccade) terminated.

The partial aspect of the partially independent pathways must be emphasized, however, as evidence for interactions between the two pathways are also present in our data. For example, although head movements are not prevented during stimulation delivered before gaze onset, they are significantly delayed (35/48 datasets). There is also a small effect on head amplitude when stimulation is delivered prior to gaze onset (significant for 21/48 datasets) and even when stimulation is triggered on gaze onset (significant for 29/67 datasets). Moreover, stimulation modified both the initial and peak velocity and acceleration of the head movements in a majority of the datasets. But note that for some datasets, the head kinematics and/or dynamics are greater in the stimulation condition compared with control trials. This effect is not likely specific to a stimulation site or to the target configuration because the enhancement was observed on only some trials within individual datasets (see Fig. 5B, for example). Finally, the ongoing head movement reaccelerates in association with the gaze shift and eye saccade that follows stimulation offset (Fig. 1) (also see Corneil et al. 2002). Evidence for interactions between the two pathways comes for other studies as well. For instance, natural or stimulation-induced variability in an ongoing head-movement dynamics alters the dynamics of the saccadic eye movement (Freedman and Quesy 2004; Freedman and Sparks 2000).

Biomechanical factors (see Tables 1 and 2), such initial head position, velocity of ongoing head movements, and inertia could also have contributed to the variability observed in the head dynamics. Despite the variability in the head-movement dynamics after OPN stimulation, there was no obvious sign of a sharp deceleration of the head movement. If the OPN stimulation had activated fibers projecting to the antagonistic neck muscles, which are lengthening during an ongoing head movement, then the force produced by the recruitment of lengthening muscle fibers would have been magnified. This would have produced a sharp and observable deceleration, and perhaps even halted the head movement. In our datasets, the deceleration was modest at best, suggesting that stimulation of the OPNs did not result in active breaking of the head movement.

Based on these results, we conclude that stimulation of the OPN region gates the neural pathway required to produce the saccadic eye component of the gaze shift and that this pathway

### Table 1. P values obtained from 2-way ANOVA on analyzed parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Stimulation Before Gaze Onset</th>
<th></th>
<th>Stimulation After Gaze Onset</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>IHP</td>
<td>Type</td>
<td>IHP*Type</td>
</tr>
<tr>
<td>Gaze latency (Fig. 2, A and D)</td>
<td>0.2118</td>
<td>0</td>
<td>0.1394</td>
</tr>
<tr>
<td>Head latency (Fig. 2, B and E)</td>
<td>0.0039</td>
<td>3.3277E-08</td>
<td>0.0852</td>
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<tr>
<td>Head re gaze onset (Fig. 2, C and F)</td>
<td>1.6245E-05</td>
<td>8.3250E-12</td>
<td>0.2811</td>
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<tr>
<td>Gaze displacement (Fig. 3B and 4B)</td>
<td>0.2775</td>
<td>8.4377E-15</td>
<td>0.3924</td>
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<tr>
<td>Head displacement (Fig. 3B and 4B)</td>
<td>0.0007</td>
<td>0.0575</td>
<td>0.4877</td>
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<tr>
<td>Eye displacement (Fig. 3B and 4B)</td>
<td>0.6758</td>
<td>0</td>
<td>0.4453</td>
</tr>
<tr>
<td>Peak velocity (Fig. 7, A and C)</td>
<td>1.5157E-06</td>
<td>0.0038</td>
<td>0.2439</td>
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<tr>
<td>Peak acceleration (Fig. 7, B and D)</td>
<td>3.8370E-05</td>
<td>0.0013</td>
<td>0.1779</td>
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<tr>
<td>Time of peak velocity (Fig. 8, A and C)</td>
<td>0.0693</td>
<td>0.0003</td>
<td>0.1679</td>
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<tr>
<td>Time of peak acceleration (Fig. 8, B and D)</td>
<td>0.0002</td>
<td>4.5796E-05</td>
<td>0.5252</td>
</tr>
</tbody>
</table>

The independent variables were initial head position (IHP: centered or contraversive to gaze shift direction) and trial type (control or stimulation).

### Table 2. P values obtained from 2-way ANOVA on analyzed parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Stimulation Before Gaze Onset</th>
<th></th>
<th>Stimulation After Gaze Onset</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ΔG&lt;sub&gt;d&lt;/sub&gt;</td>
<td>Type</td>
<td>ΔG&lt;sub&gt;d+Type&lt;/sub&gt;</td>
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<tr>
<td>Gaze latency (Fig. 2, A and D)</td>
<td>0.5291</td>
<td>1.8874E-15</td>
<td>0.9566</td>
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<td>Head latency (Fig. 2, B and E)</td>
<td>0.7672</td>
<td>1.7405E-06</td>
<td>0.6005</td>
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<tr>
<td>Head re gaze onset (Fig. 2, C and F)</td>
<td>0.0473</td>
<td>1.1102E-16</td>
<td>0.2212</td>
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<tr>
<td>Gaze displacement (Fig. 3B and 4B)</td>
<td>0.2827</td>
<td>0</td>
<td>0.3573</td>
</tr>
<tr>
<td>Head displacement (Fig. 3B and 4B)</td>
<td>0.0046</td>
<td>0.0162</td>
<td>0.3378</td>
</tr>
<tr>
<td>Eye displacement (Fig. 3B and 4B)</td>
<td>0.9055</td>
<td>0</td>
<td>0.4768</td>
</tr>
<tr>
<td>Peak velocity (Fig. 7, A and C)</td>
<td>0.0111</td>
<td>0.0002</td>
<td>0.3454</td>
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<tr>
<td>Peak acceleration (Fig. 7, B and D)</td>
<td>0.0975</td>
<td>0.0058</td>
<td>0.6556</td>
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<tr>
<td>Time of peak velocity (Fig. 8, A and C)</td>
<td>0.0070</td>
<td>1.1975E-06</td>
<td>0.1718</td>
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<tr>
<td>Time of peak acceleration (Fig. 8, B and D)</td>
<td>0.0002</td>
<td>3.4733E-08</td>
<td>0.0817</td>
</tr>
</tbody>
</table>

The independent variables were desired gaze amplitude (ΔG<sub>d</sub>: 40 or 60° desired gaze shift displacement; all movements were initiated from the same initial head position: 20° contraversive to gaze shift direction) and trial type (control or stimulation).
is not the only one involved in generating the head component of the gaze shift. This interpretation is in agreement with existing hypotheses of neural control of gaze shifts (Corneil et al. 2002; Freedman 2001; Goossens and Van Opstal 1997; Grantyn and Berthoz 1987; Guittton et al. 1990; Phillips et al. 1995). Figure 10 illustrates a schematic of the pathways and neural commands hypothesized to produce coordinated eye-head movements. It emphasizes that the pathway generating the eye component relies on a desired gaze-displacement command (\(\Delta G_d\)) and is gated by the OPNs. The drive to the neck muscles, in contrast, is a combination of a pure head-movement command (\(H_c\); pathway 1) and some formulation of \(\Delta G_d\). Evidence supporting the notion of a head-movement command, not necessarily associated with a gaze shift, comes from neural recordings in the superior colliculus (Gandhi and Walton 2006), central mesencephalic reticular formation (Pathmanathan et al. 2006a,b), microstimulation of the frontal and supplementary eye fields (Chen 2006; Chen and Walton 2005), and adaptation of eye-head coordination in the context of a limited visual field (Constantin et al. 2004). Neural signals encoding a desired gaze-displacement command have been proposed to exist at the level of the superior colliculus (Freedman and Sparks 1997a; Freedman et al. 1996; Klier et al. 2001; Munoz et al. 1991), frontal eye field (Guittton and Mandl 1978; Knight and Fuchs 2007; Tu and Keating 2000; but see Chen 2006 for an alternative hypothesis), and supplementary eye field (Chen and Walton 2005; Martinez-Trujillo et al. 2003). The \(\Delta G_d\) command, perhaps after some processing that incorporates the effect of the eye position in the orbits and the location of the stimulus (Freedman 2001), adds to the neck muscle drive at the level of the pontine burst generator (BG).

Some studies have suggested that the gaze command drive is parsed into individual, but not necessarily independent, anatomical pathways encoding eye and head movements before BG (pathway 2, Fig. 10) (Freedman 2001; Freedman and Quessy 2004; Phillips et al. 1995), whereas others have suggested that the separation occurs after BG (pathway 3) (Corneil et al. 2002; Galiana and Guittton 1992; Goossens and Van Opstal 1997; Grantyn et al. 1987; Guittton et al. 1990). These latter two options need not be mutually exclusive. While anatomical support for tectospinal and tectoreticulospinal projections have been reported in cats (Grantyn and Berthoz 1988; Grantyn et al. 1987, 1992; Isa and Naito 1995) and monkeys (Cowie et al. 1994; May and Porter 1992; Scudder et al. 1996a,b), it is not known whether they are subject to OPN inhibition.

The interpretations of our OPN stimulation results on the neck muscle drive pathways are limited because of at least two unknown factors: the functional importance of the \(H_c\) pathway during gaze shifts, and the effects of OPN stimulation on the \(H_c\) and \(\Delta G_d\) signals. If the \(H_c\) contribution is significant, then our results cannot distinguish between pathways 2 and 3. The occasional observation that head velocity is higher during the stimulation-induced interruption (Fig. 7) implicates a functionally significant role for the \(H_c\) command. On the other hand, if the \(H_c\) contribution is small or negligible, then our results—in particular, the observation that OPN stimulation does not prevent head movements—provide strong support for a drive to the neck muscles that is not gated by the OPNs (pathway 2).

Pathway 3 also remains viable because numerous head-movement characteristics of some datasets were altered, although this effect could also be explained by the effects of OPN stimulation on the \(\Delta G_d\) command. When head-restrained saccades are interrupted in mid-flight by stimulation of the OPN region, the high-frequency activity in colliculus neurons, presumably those encoding \(\Delta G_d\), is suppressed immediately (Keller and Edelman 1994; Keller et al. 2000). The stereotypical saccade-related burst neuron remains quiescent during most of the interruption period and, depending on the interruption duration, the same or another at a more rostral site discharges a high-frequency burst just before the resumed saccade. In another subset of neurons, presumably the build-up type, the low-frequency activity resumes after the initial suppression and maintains a low discharge rate during the interrupted period. Given the colliculus’ role in gaze control, it seems safe to assume that these neurons behave similarly for OPNs stimulation during head-unrestrained gaze shifts and head-restrained saccades. Accordingly, we speculate that for stimulation of the OPN region before gaze (or saccade) onset, a subset of collicular neurons will exhibit a low firing rate but that the level of activity may be attenuated and its onset may be delayed. If this activity represents an attenuated and delayed neuromuscular drive to the neck muscles, the resulting head movement is likely to exhibit a longer onset time and slower

**Fig. 10.** Simplified schematic of the flow of neural signals involved in generating coordinated eye-head movements. Two partially independent pathways provide input signals. A head-movement command (\(H_c\)) provides a drive to the neck muscles (pathway 1), but the functional importance of this pathway during gaze shifts has yet to be determined. A desired gaze-displacement command (\(\Delta G_d\)) contributes to producing the saccadic eye and head components of the gaze shift. One possibility (pathway 2) is that \(\Delta G_d\) is dissociated into separate eye and head commands before the burst generator (BG), which in turn provides a drive to only the extraocular motoneurons (\(M_N_e\)). Thus no subset of the \(\Delta G_d\) drive to the neck muscles is gated by the OPNs. Another scenario (pathway 3) is that the separation of \(\Delta G_d\) into separate eye and head pathways occurs after the burst generator elements. Note that pathways 2 and 3 need not be mutually exclusive. \(M_N_n\), neck motoneurons.
dynamics (Fig. 2B). Similarly, when OPN stimulation interrupts ongoing gaze shifts, the immediate suppression followed by low-frequency activity is interpreted as a weaker drive to the neck muscles, resulting in a slower but still continuing head movement. Following stimulation offset, superior colliculus neurons produce a high-frequency burst to generate a gaze shift. This activity also contributes to the neck muscle drive via either pathway (2 or 3) and re-accelerates the ongoing head movement.

**Feedback mechanisms**

In most models of the saccadic system, feedback control of head-restrained saccades is maintained by subtracting an internal feedback signal representing current eye displacement from a motor command of desired eye displacement (Becker et al. 1981; Robinson 1975). The resulting motor error signal drives the burst generator until feedback reduces it to zero. When the head is unrestrained, appropriate feedback analyses need to consider three signals—gaze, head, and eye in head. One class of models proposes that the feedback mechanisms preserve gaze accuracy (see review by Guitton et al. 2003). This is accomplished by computing a gaze motor error signal as the difference between desired gaze-displacement and feedback signals of instantaneous eye-in-head and head displacements (Galiana and Guitton 1992; Guitton et al. 1990). Note that the exact eye and head contributions to the gaze shift are not controlled in this schema. Another family of models suggests that the desired gaze displacement is decomposed into appropriate desired head amplitude and desired eye saccade amplitude, and the feedback mechanism serves the eye pathway only (Freedman 2001; Freedman and Quessy 2004). Gaze and head amplitudes are not controlled in this framework. These competing hypotheses have been tested by observing behavioral and electrophysiological compensation to passive head perturbations (Choi and Guitton 2006; Coimbra et al. 2000; Matsuo et al. 2004; Sylvestre and Cullen 2006), microstimulation of numerous brain structures (Freedman and Quessy 2004; Péลisson et al. 1995), and naturally prolonged movements (Bergeron and Guitton 2000, 2002). The majority of studies have demonstrated gaze accuracy (see Freedman and Quessy 2004 for exception). We induced perturbations by stimulation of the OPN region and also found that the mean difference in total gaze amplitude between the control and stimulation conditions was on the order of 1°. Thus our data are not inconsistent with the gaze comparator class of models.

To evaluate the eye comparator class of models, we could compare the total amplitude of the saccadic eye components in the control and stimulation conditions. This comparison would be justified if we can verify that the desired saccadic eye component is not recomputed during the interruption, and extrapolation of existing electrophysiological experiments on the head-restrained system suggests that this condition may be violated. Referring to the effects of interrupted saccades on SC activity discussed in the preceding text (Keller and Edelman 1994; Keller et al. 2000), the locus of activity on the SC shifts to a rostral site encoding the smaller resolicited saccade when interruption durations are long (>70 ms; Keller, personal communications), which was typical of our data. Assuming that this observation also holds for the head-restrained condition, the locus of activity would also shift to a more rostral site for the resumed gaze shift. A new desired gaze-displacement command could result in recalculating the desired saccadic component, thereby confounding interpretations of a comparison of the total saccade amplitude in the control and stimulation conditions. Hence this analysis is not reported, and we are unable to test the eye comparator class of models in a straight-forward manner.

**Summary**

We found that stimulation of the OPN region induces dissociation between the eye and head components of gaze shifts. The ocular saccade is completely inhibited by the stimulation. A head movement can be initiated during the stimulation, and an ongoing head movement continues along its trajectory. Post stimulation offset, the gaze shift resumes as a coordinated eye-head movement and preserves gaze accuracy. A reacceleration of the head movement is typically observed in association with the resumed gaze shift.

Compared with the control condition, head-movement dynamics are altered during the stimulation—typically they are attenuated but sometimes enhanced. These results indicate that the signals producing the saccadic eye-movement component of a coordinated eye-head gaze shift are not the only commands involved in the generation of the head component of the gaze shift. Additional drives to the neck muscles can stem for the desired gaze-displacement command and head-movement commands not necessarily associated with gaze shifts. This latter signal could be relayed through other volitionally recruited pathways projecting to the spinal cord and could account for an increase in head dynamics during the stimulation.

**Acknowledgments**

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**References**


