Direction-dependent excitatory and inhibitory ocular vestibular-evoked myogenic potentials (oVEMPs) produced by oppositely directed accelerations along the midsagittal axis of the head

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Abstract Oppositely directed displacements of the head need oppositely directed vestibulo-ocular reflexes (VOR), i.e. compensatory responses. Ocular vestibular-evoked myogenic potentials (oVEMPs) mainly reflect the synchronous extraocular muscle activity involved in the process of generating the VOR. The oVEMPs recorded beneath the eyes when looking up represent electro-myographic responses mainly of the inferior oblique muscle. We aimed: (1) to study the properties of these responses as they were produced by head acceleration impulses to the forehead and to the back of the head; (2) to investigate the relationships between these responses and the 3-D linear head accelerations that might reflect the true stimulus that acts on the vestibular hair cells. We produced backward- and forward-directed acceleration stimuli in four conditions (positive and negative head acceleration impulses to the hairline and to the inion) in 16 normal subjects. The oVEMPs produced by backward- and forward-directed accelerations of the head showed consistent differences. They were opposite in the phase. The responses produced by backward accelerations of the head began with an initial negativity, n11; conversely, those produced by accelerations directed forward showed initially a positive response, p11. There was a high inter-subject correlation of head accelerations along the head anteroposterior and transverse axes, but almost no correlation of accelerations along the vertical axis of the head. We concluded that backward-directed head accelerations produced an initial excitatory response, and forward-directed accelerations of the head were accompanied by an initial inhibitory response. These responses showed dependence on acceleration direction in the horizontal plane of the head. This could be consistent with activation of the utricle.

Keywords Vestibulo-ocular reflex · Otolith · Utricle · Vestibular-evoked potentials · Head acceleration

Introduction

The vestibular system senses head displacements and, in accord with other sensory and motor systems, generates appropriate reflex responses to maintain balance of the body and position of the eyes in space (Baloh and Honrubia 2001). The main role of the vestibulo-ocular reflex (VOR) is to preserve clear vision especially during rapid head displacements (Angelaki and Dickman 2000; Highstein et al. 2005). This objective is achieved by the semicircular canals and otolithic organs and their projections to the ocular motor nuclei in the brainstem (Leigh and Zee 2006). The canals sense angular accelerations, the saccular afferents are predominantly activated by linear forces acting in the vertical plane, while the utricle responds mainly to linear accelerations in the horizontal plane (Goldberg and Fernandez 1971; Fernandez and Goldberg 1976; Raphan and Cohen...
Several studies have recently investigated the electrical potentials that can be recorded close to the eyes in response to bone-conducted (BC) sound (Jombik and Bahyl 2005a; Iwasaki et al. 2007) and air-conducted (AC) sound (Halмагyи et al. 2003; Jombik and Bahyl 2005b; Aw et al. 2006; Chihara et al. 2007) as well as galvanic stimuli (Rosengren et al. 2009a, b). It is now generally accepted that these potentials mainly reflect the synchronous extraocular muscle activity that is involved in the process of generating the VOR (Rosengren et al. 2005; Todd et al. 2007). Nowadays, they are termed “ocular vestibular-evoked myogenic potentials” (oVEMPs) (Todd et al. 2007). It is probable that many, if not all, eye muscles are activated by the above-mentioned stimuli. However, the most reproducible responses can be obtained by bipolar recording from sites beneath the eyes during upward gaze. In this instance, the inferior oblique muscle gets closest to the recording site, and thus the response reflects the activity of mainly this muscle. It has been shown that AC and BC oVEMPs can be valuable tools for clinical investigation of the vestibulo-ocular reflex circuits (Iwasaki et al. 2007; Rosengren et al. 2007, 2008; Welgampola 2008; Welgampola et al. 2008; Iwasaki et al. 2008b, 2009; Wang et al. 2010; Cheng et al. 2009; Manzari et al. 2010). However, the specificity of the primary vestibular afferents and their efferent projections that are activated by different kinds of stimuli, i.e. AC and BC sound, is still a matter of dispute. In our previous study using EOG montage, we showed a complementary mirror image of the responses with phase reversal of the recorded potentials that were produced by head taps to the mastoids, i.e. in response to acceleration along the head’s transverse axis (Jombik and Bahyl 2005a). This finding was later confirmed by the oVEMP study of (Todd et al. 2008a). It is also in agreement with findings on the cervical-evoked myogenic potentials (cVEMPs) produced by lateral head taps (Brantberg and Tribukait 2002; Brantberg et al. 2009) and lateral impulse stimuli (Rosengren et al. 2009a, b). These findings might suggest utricular origin of the responses. Similar behaviour of the cVEMPs, and the periocular potentials produced by BC sound stimuli in the sagittal plane and recorded by EOG montage, has been shown in (Brantberg et al. 2008; Jombik et al. 2008). On the other hand, Iwasaki et al. (2008a) found no difference between oVEMPs produced by head taps to the hairline and inion. In another study (Lin et al. 2010), these responses differed in the latency of their first negative peak. The authors explained this finding by the differing conduction velocities of BC sound when stimuli were applied to the anterior sites of the head compared to the inion. The BC oVEMPs produced by stimuli applied to the midline of the head allow a simple and straightforward assessment of the symmetry of the vestibular function. Several studies have shown the reliability and efficacy of this method in a clinical setting (Iwasaki et al. 2008b, 2009; Wang et al. 2010; Manzari et al. 2010).

In the current study, we attempt to investigate the differences between the oVEMPs produced by BC stimuli to the hairline versus the inion and the relationship of these responses to head accelerations recorded close to the otic capsule at the mastoid. We also want to clarify some of the above-mentioned controversies regarding the BC oVEMP produced at midsagittal cranial sites.

**Methods**

**Subjects**

Sixteen healthy volunteers (mean age 38.3 years; SD 13.5) participated in the study. That included eight women (mean age 36.1 years; SD 12.1) and eight men (mean age 41.5 years; SD 14.8). All participants gave informed consent in line with the Helsinki Declaration, and the study was approved by the local ethics committee.

**Experimental procedure**

The experiment consisted of two parts. In the first part, EMG responses of the oVEMPs were produced and recorded. The other part consisted of the recording of head accelerations.

**Stimulation**

The stimuli consisted of positive and negative impulses. The driving electrical waveforms for the impulses (positive or negative square waves with duration of 1 ms) were delivered by the sound generator of a 5-channel EMG/EP system (Teca Synergy, Medelec, UK), and the same apparatus was used for recording the EMG signals and the head accelerations. The impulses were amplified and low-pass filtered at 250 Hz by an external amplifier and applied using a custom-made mini-shaker at a rate of 3 per second and 1 per second for producing oVEMP and recording head accelerations, respectively. The mini-shaker was fitted with a short 2-cm bolt terminating in a bakelite cap 1.5 cm in diameter, which was the contact point for the stimulator on the subject’s head. The stimulator was held normal to the subject’s head with an approximate force of 10–20 N by the experimenter, who stood behind or in front of the subject according to the site of stimulation. Owing to the electromechanical properties of the mini-shaker, the filtering and the stimulus application mode, the main frequency of the resultant output impulse was about 100 Hz. The resultant output represented the impulse response of the coupled shaker-head system. The net effect was the acceleration of
this coupled shaker-head system that was directed alternately away from and towards the experimenter by the positive and the negative impulses, respectively.

OVEMP recordings

EMG signals were recorded by bipolar montage. The active (inverting) electrode was placed on the orbital margin below the eye and referred to an electrode approximately 20 mm below it on the cheek, and the ground electrode was placed on the sternum. The EMG signals were amplified and band-pass filtered (5–500 Hz), and sampled at 20 kHz for 50 ms following stimulus onset. During stimulation, the subjects were seated comfortably and instructed to look up and relax their jaw muscles. EMG responses to 30 stimuli were averaged for each of the four conditions, i.e. positive and negative impulses that were subsequently applied on the subject’s head at Fz1 (midline forehead at hairline) and inion.

Accelerometry

In the second part of the experiment, linear acceleration of the head was recorded using a custom-made miniature tri-axial accelerometer consisting of two ADXL311 dual-axis analogue devices cemented together and embedded in synthetic resin. Another dual-axis accelerometer of the same type was fitted to the bottom of a cylindrical plastic nozzle that now covered the baclelite cap of the stimulator. Thus, the active axis of the accelerometer, which was oriented normal to the head, directly recorded the acceleration impulse at the contact sites of the stimulator and the head, because the acceleration pulse was transmitted through the accelerometer. Linear acceleration of the head was produced by the same stimuli used for producing EMG responses and was recorded at the left mastoid bone site approximately 1 cm behind the pinna of the ear at the level of the tragus. To ensure tight contact of the accelerometer with the head surface, the subjects pressed the miniature device to their heads with their index fingers. The placement of the accelerometer was constant in all recordings, and its axes were oriented in parallel with the anteroposterior X, the vertical Z and the transversal Y axes of the head. The 3-D accelerometer signal was amplified and band-pass filtered (0.2–500 Hz), and sampled at 20 kHz for 50 ms following stimulus onset. Five to ten head acceleration stimuli were averaged for each condition.

Auxiliary experiments

The applied impulses could cause not only translational but also rotational displacement of the head; thus, they could stimulate not only the afferents of the otolithic organs but also the canal afferents. In order to pursue this possibility, the experimental procedure was repeated in a fixed head condition in one subject. The subject (one of the authors, PJ) pressed his head to the wall on the side opposite to the site of stimulation.

Data analysis

Latencies and amplitudes were measured for both the EMG and accelerometer signals off line. The latencies were analysed both for their absolute values and for their values corrected to the mini-shaker acceleration onsets. Absolute amplitudes were measured at the response peaks, and relative amplitudes were calculated as the differences between the subsequent oppositely directed peaks. Repeated measures ANOVAs and t tests when appropriate were used to compare response amplitudes and latencies. Reproducibility was assessed with Pearson’s correlations. Frequency analysis of the stimulus and accelerometry records at the mastoid and of the EMG responses was performed on grand averages of the traces using FFT. Values reported are means ± SD. For conciseness, only the results of the acceleration onsets and the first acceleration peaks recorded at the mini-shaker and at the mastoid site and the initial two peaks of the VEMPs are displayed here.

Results

Accelerometry measurements on the mini-shaker

Owing to the electro-mechanical delay of the hardware, the onset of mini-shaker acceleration began at about 0.50 ms after the trigger. It reached its peak value of about 1.3 g at 2.20 ms (Table 1). There were no statistically significant differences in the acceleration latency and magnitude at this first peak between the four experimental conditions. However, there was a small but statistically significant difference in the mini-shaker acceleration onset between stimuli applied to the hairline at Fz and to the inion (P < 0.01). Latency was about 30 µs shorter in stimuli applied to the inion compared to stimulation at Fz. On the other hand, there were no significant differences between the latencies of the first peak of mini-shaker acceleration.

Accelerometry measurements at the mastoid. Comparison of the findings on individual (x, y and z) axes between the four experimental conditions

Onset of acceleration along all 3 axes of the accelerometer at the mastoid was between 1.5 and 2.5 ms, while the peak
values of about 0.02–0.06 g were reached between 3.5 and 5.5 ms (Table 2). There were no significant differences either between the latencies of acceleration onsets or the latencies of first peaks along the head X axis. On the other hand, the latencies of the onsets and the first peaks along the Y axis as well as the latencies of the first peaks along the Z axis were significantly shorter for stimuli applied to Fz ($P < 0.05$). The magnitudes of the first acceleration peaks along all the axes in the four experimental conditions did not show significant differences.

However, the orientations of the acceleration vectors along the head X and Y axes showed consistent differences that were dependent on the initial direction of the stimulus delivered by the mini-shaker, i.e. the direction of mini-shaker acceleration. Positive impulses applied to Fz and negative impulses at the inion caused backward-directed head acceleration along the head X axis and outward (laterally)-directed acceleration along the head Y axis (Fig. 1). In contrast, positive impulses applied to the inion and negative impulses applied to Fz resulted in forward-directed acceleration along the head X axis and inward (medially)-directed acceleration along the Y axis (Fig. 2). Thus, the acceleration vectors along the head X and Y axes were out of phase with forward- and backward-directed stimuli.

The acceleration directions along the head X axis were identical in all test subjects, and the mean values of inter-individual Pearson’s correlation coefficients “r” of the acceleration traces during the first 6 ms showed very high correlations (+Fz 0.84, +Inion 0.93, —Inion 0.90, and —Fz 0.80).

In one subject, the acceleration vectors along the head Y axis were inverse compared to the rest of the test subjects for stimulation at Fz. Another two subjects were inverse to the rest of the subjects for stimuli applied to the inion. All these three “inverse” subjects were men. Despite these few exceptions, the mean values of the inter-subject correlation

Table 1 Accelerometry findings at the interface of the mini-shaker and the head

| Acceleration backward | Acceleration forward |
|-----------------------|----------------------|
| Fz plus | Inion minus | Inion plus | Fz minus |
| Ons lat (ms) | Mean 0.51 | 0.48** | 0.48** | 0.51 |
| | SD 0.06 | 0.06 | 0.04 | 0.05 |
| Peak 1 lat (ms) | Mean 2.16 | 2.21 | 2.18 | 2.18 |
| | SD 0.05 | 0.06 | 0.04 | 0.07 |
| Ons-peak 1 amp (g) | Mean 1.22 | 1.31 | 1.27 | 1.23 |
| | SD 0.20 | 0.14 | 0.14 | 0.19 |

Ons onset, Lat latency, g = 9.81 m s$^{-2}$

** Statistical significance level at $P < 0.01$ for shorter latencies of the acceleration onsets in stimuli applied to the inion

Table 2 Accelerometry findings at the mastoid site

| Acceleration backward | Acceleration forward |
|-----------------------|----------------------|
| Fz plus | Inion minus | Inion plus | Fz minus |
| X | Y | Z | X | Y | Z |
| Ons (ms) | Mean 2.09** | 1.53$^\#$ | 2.18** | 2.14** | 1.69 | 2.09 |
| | SD 0.54 | 0.41 | 0.43 | 0.39 | 0.26 | 0.78 |
| Peak 1 lat (ms) | Mean 4.99**$^\#$ | 3.98$^\#$ | 4.44$^{**\#}$ | 4.86** | 4.11 | 4.89** |
| | SD 0.46 | 0.26 | 0.73 | 0.56 | 0.56 | 0.61 |
| Ons-peak 1 amp (g) | Mean 0.06$^{**\#}$ | 0.05 | 0.03 | 0.05$^{**\#}$ | 0.04 | 0.03 |
| | SD 0.03 | 0.03 | 0.02 | 0.02 | 0.02 | 0.01 |

Ons onset, Lat latency, Sh mini-shaker, g = 9.81 m s$^{-2}$

$^\#$ and $^{**\#}$ statistical significance level at $P < 0.05$ and $P < 0.01$ for shorter latencies of the acceleration onsets and the first peaks on the Y and Z axes for stimuli applied to Fz in the between-conditions comparisons

$^*$ and $^{**}$ statistical significance level at $P < 0.05$ and $P < 0.01$, respectively, for longer latencies of the acceleration onsets and the first peaks on the X and Z axes compared to the Y axis in the within-conditions comparisons

† and $^{**\#}$ statistical significance level at $P < 0.05$ and $P < 0.01$, respectively, for longer latencies of the acceleration first peaks on the X axis compared to the Z axis in the within-conditions comparisons

$^\#$ and $^{**\#}$ statistical significance level at $P < 0.05$ and $P < 0.01$, respectively, for higher amplitudes of the first acceleration peaks on the X and the Y axes compared to the Z axis in the within-conditions comparisons

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coefficients were still moderately high (+Fz 0.65, +Inion 0.46, −Inion 0.46, −Fz 0.62).

On the other hand, the acceleration vectors along the head vertical Z axis showed substantial variations between test subjects in all four experimental conditions, and the mean values of inter-individual correlation coefficients of Z acceleration traces during the same time interval were low or almost negligible (+Fz 0.05, +Inion 0.23, −Inion 0.36, −Fz 0.03). However, more detailed analysis revealed that from stimuli producing backward-directed head acceleration, i.e. +Fz and −Inion, the test subjects displayed two patterns of acceleration along the Z axis and three patterns from stimuli producing forward-directed head acceleration, i.e. +Inion and −Fz. In general, more subjects showed an upward-directed acceleration vector for backward-directed stimuli and vice versa. Nevertheless, this entirely different out-of-phase behaviour of the acceleration vectors along the head Z axis was not reflected in any parameters of the oVEMP responses as shown in Figs. 3 and 4.

Accelerometry measurements at the mastoid. Comparison of the findings between individual (x, y and z) axes within the four experimental conditions

In all four experimental conditions, the latencies of the onsets and the first peaks were shortest for accelerations along the head Y axis (Table 2). These differences between the Y and X axes were statistically significant in all four experimental conditions, both for the latencies’ absolute values and for the values corrected to the mini-shaker acceleration onsets (P < 0.01). These differences were also significant between the Y and Z axes (P < 0.05) in all four experimental conditions, except for the onset by negative stimuli applied to the inion. Between the X and Z axes, there were no significant differences in the latencies except for the first peak produced by stimuli to Fz where Z acceleration peaks occurred significantly earlier (P < 0.05). The acceleration magnitude at the first peak tended to be highest for the X axis and lowest for the Z axis,
Fig. 2 EMG responses and accelerations of the head produced by stimuli directed forwards.

Fig. 3 Relationship of the EMG responses and accelerations of the head along the vertical axis produced by stimuli directed backwards. The two panels at the top show the responses produced by positive impulses at the hairline at Fz, and the panels at the bottom display the responses to negative impulses at the inion. The panels on the left side show the grand average (bold) traces and responses in the individual subjects superimposed. For better clarity, only the grand average traces are displayed in the panels on the right side. In each panel, the upper two rows show the EMG responses and the bottom two the acceleration traces of the same subgroups. Both stimuli produced an initial upward-directed acceleration of the head in the majority of subjects. However, by positive stimuli to the hairline and by negative impulses to the inion, the acceleration of the head was opposite in five and two subjects, respectively. The oppositely directed vertical accelerations did not show any substantial influence on the EMG.
and this difference was significant \((P < 0.01)\). There were no significant differences between the \(X\) and the \(Y\) axes. Between the \(Y\) and the \(Z\) axis, the difference was significant only for positive impulses applied to the inion \((P < 0.05)\).

### oVEMPs

In general, there were no significant differences in the latencies and amplitudes between the oVEMP responses recorded from the left and right eye. Asymmetry ratios of peak-to-peak amplitudes that were calculated using the standard Jongkees formula\(^2\) were well below 34 per cent. We therefore used the mean values from both eyes for subsequent analyses. However, there were a few exceptions when some subjects in one of the four conditions showed asymmetrical responses with low amplitude or distortion on one side. In such cases, the better response was used for analysis.

In all four conditions, the oVEMP responses showed consistent behaviour, and they copied the shapes of \(X\) acceleration traces with a delay of about 5–6 ms, as shown in Figs. 1 and 2. The responses that were produced by backward-directed acceleration of the head showed an initial negative peak at about 11 ms. In contrast, the responses produced by forward-directed accelerations began with an initial positivity with approximately the same latency (Table 3). The subsequent peaks that occurred

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\(^2\) Asymmetry ratio \( [%] = \frac{[\text{larger peak amplitude} - \text{smaller peak amplitude}]}{[\text{larger peak amplitude} + \text{smaller peak amplitude}]} \times 100.\)
at about 14, 19, 24, 30 and 38 ms showed similar out-of-phase behaviour (Fig. 5). Only in two subjects, both women, were there some latency shifts in inion responses. They partially disrupted the otherwise perfect appearance of the out-of-phase behaviour of the responses that were still preserved in responses produced by stimuli with opposite polarity to Fz in these two subjects.

There were no significant differences either between latencies of the onsets or latencies of later-occurring subsequent peaks of the oVEMP responses in the four experimental conditions except for the second peak that occurred at about 14 ms. Both the absolute latencies of this peak and its latency adjusted to the mini-shaker acceleration onset were significantly shorter for stimuli producing forward acceleration, i.e. positive impulses to the inion and negative impulses to Fz ($P < 0.05$). However, there were no significant latency differences for this peak between conditions with the same acceleration direction as shown in Table 3.

The amplitude of the first oVEMP peak was significantly highest ($P < 0.01$) for positive stimuli to Fz in comparison with the other three experimental conditions, but there were no significant differences between the rest of the conditions. Amplitudes measured between the first and the second peak were higher for stimuli applied to Fz. These differences were significant ($P < 0.01$) except for $+$Inion versus $-$Fz as shown in Table 4.

Comparisons of the accelerometry and oVEMP

in head-free and head-fixed conditions

Accelerations of the head and the oVEMP responses tested in one of the subjects in all four experimental conditions were almost entirely identical on visual assessment regardless of whether the head was free or fixed (Fig. 6).

Frequency analysis

Frequency analysis of the mean traces of the stimulus recorded at the contact site of the head and the mini-shaker and at the site of the mastoid, as well as the EMG (oVEMP) signal, showed that the power was approximately 100 Hz (the exact values were 117 for the mini-shaker; and 78 Hz for X and Z axes as well as EMG and 98 Hz for Y axis) as shown in Fig. 7.

Discussion

In general, we found almost consistent behaviour in head acceleration in the horizontal plane and resultant oVEMP responses in the studied sample of normal subjects. The traces of the oVEMPs that were produced by oppositely directed head accelerations consistently showed the opposite phase.

Accelerometry

Regardless of the negative or the positive stimuli, the direction of the head acceleration always followed the direction of movement of the stimulation device. Owing to the spring-like mechanics of the neck, the tight contact caused by the pressure of the mini-shaker against the head resulted in the head and the device moving in synchrony (the elastic restoring forces of the skull might also play some role). The

Table 4  oVEMP amplitude findings

| Acceleration backward | Acceleration forward |
|-----------------------|----------------------|
| Fz plus               | Inion minus          | Inion plus | Fz minus |
| n11/p11 (µV)          |                      |            |          |
| Mean                  | 5.21**               | 2.10       | 1.57     | 2.44     |
| SD                    | 3.05                 | 1.25       | 1.25     | 1.86     |

n11/p14/n14 (µV)

| Mean                  | 10.85*#              | 4.70       | 6.08     | 9.00#    |
| SD                    | 4.92                 | 2.33       | 2.38     | 5.36     |

For abbreviations see Table 3

** Statistical significance level at $P < 0.01$ for higher amplitudes of the first oVEMP peaks between $+$Fz and the rest of the conditions in the paired comparisons

* Statistical significance level at $P < 0.01$ for higher amplitudes of the second oVEMP peaks between $+$Fz versus $+$Inion and $-$Inion in the paired comparisons

# Statistical significance level at $P < 0.01$ for higher amplitudes of the second oVEMP peaks between $-$Fz versus $-$Inion in the paired comparisons
acceleration traces on the mini-shaker and the head displayed a damped oscillation pattern.

The latency of mini-shaker acceleration onset was about 30 μs shorter for stimuli applied to the inion compared to stimulation at Fz. The reason for the earlier onset of the mini-shaker acceleration for stimuli applied to the inion might be the more compliant hair layer at this site in contrast to the less compliant bare skin at the hairline.

The magnitudes of the acceleration peaks recorded at the site of the mastoid were about two orders below those recorded at the contact site of the mini-shaker and the head. These values are similar to those described in previous studies by other authors (Iwasaki et al. 2008a; Todd et al. 2008a; Rosengren et al. 2009a, b).

The response of the skull to BC stimuli is complex. These mechanical stimuli can produce both elastic-compression waves and in-compressive vibration of the head. The displacements of the head can consist of three-dimensional translation and rotation. The size of these displacements would have been very small, at the order of about 10⁻⁶ m, given the size and the duration of the accelerations which we have recorded on the heads (Todd et al. 2008a). At low frequencies below 1 kHz, the head and especially the skull base tends to move as a whole (Stenfelt and Håkansson 2000; Stenfelt and Goode 2005a, b). Nevertheless, the shell of the skull and its base tend to behave differently, so the elastic-compressive waves in the shell can still be prominent even for low-frequency stimuli. However, the accelerometry findings in this study showed that, despite the low frequency of the stimulus at about 100 Hz and mode of its application, i.e. pressing the mini-shaker to the head, a significant portion of the elastic-compression waves was transmitted from the skull shell to the mastoid. We suspect that the reason for this is the site of the stimulation, i.e. the shell of the skull, because despite the highest magnitude of skull acceleration occurring along its X axis, in contrast to studies that used 500-Hz BC tone burst stimuli and head taps (Iwasaki et al. 2008a, 2009), the compression waves oriented along the head Y and Z axes were still present and showed significant magnitudes.

Interpretation of the accelerometric findings is not easy, but the following could be proposed: the skull is neither a homogeneous nor an isotropic mass body. Compliance of the skull bone can vary at different sites. Moreover, its anterior part, including the frontal, sphenoid, and the temporal bones are joined together by tightly ossified sutures. By contrast, the occipital bone at the base of the skull has a wedge shape. The lateral parts of this wedge are joined with the pyramidal parts of the temporal bones at least partly by softer fibrocartilaginous tissues.

Owing to the compliance of the skull shell bone, the applied force of stimulus first caused compressive-elastic
deformation of the skull, and only after exhaustion of this elastic freedom did acceleration in accordance with the direction of the stimulus follow. This could explain the longer latencies of the acceleration along the head X axis in all four experimental conditions.

The reason for shorter latencies of acceleration along the Y and Z axes for stimulation at the Fz site compared to the inion might be the shorter transmission time for this elastic-deformation wave from the frontal shell to the mastoid bone through the hard dense bone structures in contrast to the softer connections in the path from the inion. Nevertheless, according to the latency differences found in this study, the differences in the conduction velocity could be only two- or several-fold faster, but should not reach values of the order suggested in one previous study (Lin et al. 2010).

The differences in the orientation of Y axis acceleration vectors produced by backward- and forward-directed stimuli suggest an entirely different mode of skull deformation by these oppositely directed stimuli. Owing to the tight osseous connections of the frontal and temporal bones, the initial elastic-deformation forces produced by stimuli applied to the hairline cause transient changes to the skull shape from spherical to ovoid. This leads to outward-directed acceleration of the pyramids. On the other hand, the forces acting at the inion might cause small displacements of the wedge-shaped basal part of the occipital bone with concomitant elongation and narrowing of fibrocartilaginous structures in the foramen lacerum and subsequent medially directed acceleration of the pyramids. We propose this model for the positive stimuli, and the opposite might happen with negative impulses.

Head acceleration direction in the utricular and saccular planes and the oVEMPs

The directions of the head accelerations along the anteroposterior and the transverse axes, i.e. in the main plane of the utricle, were with few exceptions the same in the tested subjects. On the other hand, there was a great variability in the acceleration along the head vertical axis, i.e. in the main saccular plane, recorded at the mastoid bone without any consistent trend between the subjects, the polarity of the stimulus, and the stimulation site. The oVEMP responses also showed consistent behaviour in their phases in relation to the head acceleration direction in the plane of the utricle, but not to the acceleration direction in the saccular plane. Thus, in general the head impulses that were initially directed backward (positive impulses to the hairline and negative impulses to the inion) also caused backward- and
outward-directed acceleration at the mastoid bone. The resultant oVEMP potentials were consistently accompanied by an initially negative response with a mean latency of about 11 ms, i.e. the n11 peak. Conversely, forward-directed accelerations of the head (negative impulses to the hairline and positive impulses to the inion) almost always caused inward-directed acceleration at the mastoid bone in addition to the forward-directed vector and produced an initially positive oVEMP response with approximately the same latency of about 11 ms but with opposite polarity, i.e. the p11 peak.

The EMG responses show initial negativity when recorded close to the motor point of the muscle and when the muscle is depolarized. Conversely, the inhibition of the ongoing muscle tonic activity is accompanied by positive potential (Colebatch and Rothwell 2004). Direct eye movement recordings have shown that the first oVEMP potential reflects the activity of the inferior oblique muscle (Welgampola et al. 2009). Thus, we might expect an excitatory response of the inferior oblique muscle to backward-directed head acceleration (and vice versa).

The oVEMP potentials differed in their phases, i.e. polarities, but otherwise they were almost identical. Nevertheless, the magnitudes of the first two peaks were higher for stimulation at the Fz site. This might reflect better efficacy of stimulation at this site in comparison with the impulses applied to the inion. On the other hand, the lower amplitudes of the first positive inhibitory potentials produced by negative impulses to Fz and positive impulses to the inion might reflect the natural propensity of the vestibular system for enhancing the excitatory responses.

In addition, the latencies of the later peaks were almost identical regardless of the stimulus site or polarity. We suppose therefore that they reflect the activity of the same vestibular end-organ and the same pathways, i.e. the projections from the utricle.

On the other hand, a smaller proportion of the hair cells situated in the anterior part of the saccule have their polarization vectors oriented along the anteroposterior axis of the head. Thus, some small contribution to the responses from this part of the saccule cannot be excluded.

Due to the limitations of the acceleration recording mode, we cannot exclude the possibility that head taps caused some angular head displacement around the transverse axis. Such stimuli could activate the vertical canal afferents. The anterior and posterior canals and the utricle project strongly towards the oblique muscles, while the horizontal canals and saccule do not (Isu et al. 2000; Uchino et al. 1996). Nevertheless, if the oVEMP potentials were produced by stimulation of the vertical canals, the responses must have opposite polarity. Clearly, head rotation backward will activate the posterior canal VOR, which is associated with the inhibition of the inferior oblique muscle (and vice versa). Finally, the auxiliary experiment with the head-free versus head-fixed conditions in one subject did not show any important difference in head acceleration direction or oVEMP response phases. Although this experiment could not entirely eliminate head rotation, we think that this condition could at least reduce it considerably.

Mechanism by which BC stimuli activate vestibular receptors and the origin of oVEMP responses produced by these stimuli applied to the hairline and to the inion

Young et al. (1977) found that both canal and otolithic primary afferents can be activated by BC vibrations. Later, Curthoys et al. (2006) showed that these stimuli selectively activate irregular otolithic receptors. The precise mechanism by which BC sound activates the otolithic receptors is not clear. With regard to hearing of BC sound, some studies propose that the dominant mechanism in the initiation of the cochlear bone conduction response is the action of pressure waves in the cerebrospinal fluid. These waves would be transmitted via the physiological third windows to the inner ear fluids (Sohmer et al. 2000; Freeman et al. 2000; Sohmer and Freeman 2004). However, such a mechanism would preclude direction-dependent vestibular responses to BC vibration. Thus, it is most probable that the otolithic receptors are activated by virtue of their inertia. It has been shown that the resonance for the oVEMP response occurs with frequencies around 100 Hz, where an extraordinary level of vestibular sensitivity to these responses has been found (Todd et al. 2008b). Both central and peripheral mechanisms might be responsible for this tuning. At the level of the sensory organs, the biomechanical properties of the saccular and utricular maculae are quite different. The ratio of saccular vs. utricular frequency resonance has been determined at 6:1 (Todd et al. 2009a). Owing to these peripheral biomechanical factors, the frequency tuning of the saccular afferents is close to 500 Hz. Conversely, the best frequency for the utricle is around 100 Hz (Todd et al. 2009a). Finally, the central VOR pathways for the oVEMP potentials, the afferent neural system from the extra-ocular muscles play a role in the reinforcement of oVEMP's tendency for resonance at 100 Hz (Todd et al. 2009b). The main frequency of the stimuli produced by our mini-shaker was about 100 Hz, as was the skull vibration response that we recorded at the mastoid bone.

The direction of the applied acceleration along the head X axis was the major determinant of the response. We conclude therefore that low-frequency vibrations of the head induced by impulses to the hairline and the inion probably activate utricular afferents with oppositely directed spatial polarization vectors. The resultant direction-dependent
excitatory and inhibitory oVEMP responses most likely reflect activity of the utricle-ocular VOR pathways.

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