Short latency responses in the averaged electro-oculogram elicited by vibrational impulse stimuli applied to the skull: could they reflect vestibulo-ocular reflex function?

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Objectives: To investigate whether vibrational impulse stimuli applied to the skull can be used to evoke the vestibulo-ocular reflex (VOR) and detect vestibular lesions.

Methods: Twenty four patients with unilateral vestibular loss (UVD), five with bilateral vestibular loss, two with ocular palsies, and 10 healthy subjects participated. Vibrations of the skull were induced with head taps and with a single period of 160 Hz tone burst on the inion, vertex, and the mastoids while the patients viewed a distant target. Several patients were also examined while viewing a near target, with eccentric gaze and in tilted postures. Responses were recorded by EOG.

Results: Responses occurred between 5 ms and 20 ms and seemed to be compensatory to the second phase of the sine wave of vibration impulse and were greatly diminished/absent in patients with bilateral VD and ocular palsies. The patients with UVD had asymmetrical responses in the vertical EOG with stimuli applied on the inion and vertex, with enhancement of the response amplitude on the side of vestibular loss and/or diminution on the healthy side. The asymmetry ratios between the healthy subjects and patients with UVD, and among patients with UVD were statistically significant. Some gaze and positional influences could be demonstrated consistent with otolithic reflexes.

Conclusion: If the asymmetric responses to skull vibration in UVD result from passive oscillatory movements of the orbital tissues they may reflect the otolith mediated sustained skew torsion. Conversely, if generated by active eye movements, their likely origin is a phasic VOR.

Both primary otolithic and canal afferents of a monkey can be activated by vibration. The lowest phase locking thresholds have been determined at −70 to −80 dB and median values in the most sensitive frequency range (200–400 Hz) at −20 to −40 dB of gravitational acceleration. It is still not clear whether activation of vestibular receptors by vibration has the same mechanical basis as the response to more physiological head movements. Mechanical factors are not the only determinants of response dynamics since vestibular nerve fibres can show a frequency dependent increase in gain greater than that predicted for the mechanics of sensory end organs. The mechanics of the otolithic membrane can be approximated by a damped second-order system with a resonant frequency of the order of 50–500 Hz. Thus, in contrast to the cupula–endolymph system in which the upper frequency limit is set below 60 Hz, the otolithic membrane is much better suited for transmission of bone vibrations in the audio frequency range. Conversely, canal neurones tend to be more irregular than otolith neurones, and hence might be expected to have lower vibration thresholds.

The primary function of the vestibulo-ocular reflexes (VOR) is to provide short latency compensatory eye movements early in the movement, before visual tracking comes into play. Since both the angular VOR (aVOR) and compensatory linear VOR (lVOR) operate with high pass characteristics relative to head velocity input, both might respond to skull vibrations. Both aVOR and lVOR are modulated by viewing distance, or more precisely by ocular vergence. However, this dependence is much stronger for lVOR. Moreover, the naso-occipital lVOR is also strongly influenced by gaze eccentricities relative to the naso-occipital axis. Latency of the human aVOR is approximately 10 ms with only slight intersubject variation (range 6–15 ms), coexistent with a three neurone arc. The default gain of the human aVOR at onset, elicited by low and modest accelerations is appropriate for distant targets. Higher accelerations activate responses without delay, for the expression of the effects produced by distance. Disynaptic connections between otoliths and oculomotor neurones have been demonstrated in cats, and studies in monkeys have found the lVOR latency to be similar to that of the aVOR. The mean latency of human lVOR is approximately 30 ms with high intersubject variability and gain dependent on the target distance from the onset. However, similar to the aVOR, the latency of the lVOR can be very brief in some subjects, and hence a disynaptic neurone arc may operate in humans as well. It has been suggested that fast projections might carry a baseline lVOR signal independent of viewing distance. Moreover, it is possible that compensatory lVOR uses pathways in common with the aVOR. The dynamics of the lVOR dramatically enhance the responses to frequencies of linear accelerations above 0.5 Hz independent of viewing distance. In contrast with compensatory lVOR, the orienting lVOR codes orientation of the eyes in space. Its purpose is to produce responses to static or low frequency tilts of the head relative to constant gravitational force. However, the oculomotor responses required to compensate tilt and translation differ. Unlike the translational compensatory lVOR, there is no geometric requirement for tilt responses to be modulated by changes in fixation distance. The orienting lVOR operates with low pass dynamics, and contrary to transient ocular movements generated by compensatory aVOR and aVOR, it
induces appropriate eye rotations and/or torsions that tend to be more sustained.15 17

Experimental and clinical data suggest that vibrational stimuli can evoke vestibulospinal and possibly also vestibulo-ocular responses.18–24 Thus short pulses of vibration may be appropriate for recording the earliest part of the VOR, provided that mechanical artefacts and other cranial reflexes do not obscure these responses.25 By applying vibrational stimuli to the skull, most of the mechanical energy dissipates at the bony interface and the acceleration transmitted to bone is attenuated by ~20 dB.26 Vibrations are then conducted through bone with only minor losses <5 dB.27 The velocity of propagation is estimated to be 260 m/s.28

In a preliminary study skull vibrations were generated by head taps or by sound stimuli conducted in bone. Head accelerations generated by stimulation along the naso-occipital (x) axis were measured by an accelerometer attached to the forehead. The maximal accelerations were about 0.55 cm/s^2 and 0.25 cm/s^2 for head taps and 160 Hz bone conducted tone bursts, respectively. The stimuli elicited transient responses with short latencies in the averaged electro-oculography (EOG) that may be compatible with a disynaptic VOR. If the recorded events were generated solely by horizontal or vertical eye rotations, the amplitudes of eye movements would approach ~1–4° in the responses elicited by head taps and 0.25–1° in the bone conducted tone bursts. The responses in vertical EOG were generally symmetrical in normal subjects. However, sometimes surprisingly asymmetrical responses were found in otherwise normal subjects with craniofacial asymmetries (CFA). The reason may be the different orientation of vestibular end organs on the left and right sides.29 Moreover, owing to asymmetry of skull thickness the sensitivity of the labyrinths to vibrations may be different. There was a tendency for diminution of the responses in older subjects, and there were no responses in cadavers.

The purpose of the current study was to investigate the utility of head taps and bone conducted short tone bursts as an effective means to test the function of the VOR.

METHODS

Patients

Two groups of patients with unilateral vestibular dysfunction (UVD) were investigated. The first group comprised 10 patients with total unilateral vestibular deafferentation due to surgery for an acoustic neuroma. There was no evidence of significant brainstem or cerebellar disorder. They were tested within five weeks to 12 years of surgery with a mean interval of 45 months. Five were also investigated before the surgery. The second, more heterogeneous, group comprised 14 patients with UVD of various causes with consistent abnormality of the vestibulo-collic reflex (VCR) elicited with head taps. Five patients with profound bilateral vestibular loss (BVD) were also investigated. Finally, a patient with unilateral third nerve palsy, and another with advanced stage of progressive external ophthalmoplegia were also tested. The control group comprised 10 normal volunteers without a history or clinical signs of vestibular or ocular motor disorder or visible CFA. A requirement for normal results of a VCR elicited with taps were added to the clinical inclusion criteria to reduce the likelihood of the bias from CFA. The study was performed in accordance with the 1964 Declaration of Helsinki protocol and with approval of the institutional ethics committee.

Test procedures

The subjects were seated upright in a dimly lit room, and viewed a small illuminated target directly ahead at a distance of 3.5 m. Several subjects were also examined viewing near targets at distances of 25 cm and/or 15 cm in eccentric horizontal and vertical gaze positions, as well as lying supine and prone and with eyes closed. Care was taken to ensure that the subjects’ jaw muscles were relaxed. Skull taps were applied approximately at a rate of one to two per second. The reflex hammer was fitted with an inertial switch that produced a delay of 2–3 ms, so the latencies of the recorded responses were apparently shortened by the same amount. Bone conducted sound stimuli consisted of single period of 160 Hz logon (which is a particular type of tone burst with a raised cosine, instead of trapezoidal enveloping function) and were delivered by a clinical bone vibrator with a repetition rate of three per second. All patients were investigated using the first method. The second method was used only in the first UVD group (that is, operated patients). Stimuli were applied on the inion, the vertex, and both the mastoids, thereby acceleration of skull was caused along three mutually orthogonal axes, the naso-occipital (x), dorsoventral (z), and interaural (y). However, due to the route of spread of the vibration waves to the labyrinths, the stimulation was really orthogonal only for the stimuli along the naso-occipital and interaural axes. Responses were recorded by means of Ag/AgCl surface electrodes producing bilateral monocular vertical and binocular horizontal EOG. The signal was amplified and band pass filtered (5–2000 Hz), and peaks before 20 ms after stimulus onset. Onsets occurred between 5 ms and 15 ms, and peaks before 20 ms (figs 1–3). These responses were biphasic or triphasic responses were recorded 5–20 ms after stimulus onset. Biphasic or triphasic responses were recorded 5–20 ms after stimulus onset. Onset responses in all EOG records of the patient with progressive external ophthalmoplegia and in two of the five patients with BVD, and were greatly diminished in the other three patients with BVD. Responses were also absent ipsilaterally in the vertical EOG records of the patient with third nerve palsy (fig 4). In horizontal EOG records elicited by tapping on the mastoids there was a small passive response with a zero latency culminating in the first 5 ms. Following the stimulus generated by the bone vibrator, an electrical artefact comprised the initial parts of the records up to 9 ms. Otherwise, regardless of the mode of elicitation the responses were similar in latency and shape.
Responses elicited by stimuli along the interaural (y) axis
In the binocular horizontal EOG channel the main deflection of the trace was rightward directed with stimuli applied to the left mastoid and vice versa. The responses of the left and right eyes in the vertical EOG channels were disconjugate. The amplitude of the response on the ipsilateral side of the stimulation was larger than on the contralateral side and the peak latencies of the left and right eyes were also different. However, responses were mirror images of each other, with a reversal in direction when comparing responses elicited from the left versus the right side (fig 1). These symmetrical relations were lost in patients with UVD, due to a decrease of the amplitude on the healthy side and direction/latency shifts (fig 2).

Responses elicited by stimuli along the naso-occipital (x) and dorsoventral (z) axes
The stimuli elicited symmetrical biphasic or triphasic responses with a dominant upwards directed component in the vertical EOG channels only. Responses were symmetrical in normal subjects, but asymmetrical with enhanced amplitudes on the paretic and/or decreased amplitudes on the healthy side in patients with UVD (fig 3). Inter-side amplitude difference was statistically significant for both subgroups of patients with UVD (fig 5). This asymmetry was
Kinematic considerations and positional effects

Responses were similar whether the eyes were closed or open. Inconsistent findings were obtained with fixation of a near target: the responses were unchanged or even showed decreased amplitudes. Gaze influences were studied in responses elicited from stimuli along the naso-occipital (x) axis. Upward gaze increased whereas downward gaze decreased the amplitude of responses in the vertical EOG channels. Horizontal gaze increased the amplitude on the side of the adducting and decreased it on the side of the abducting eye in healthy subjects as well as patient with UVD. Stimulation during horizontal gaze deviations also generated reverse directed responses in the horizontal EOG channel—that is, responses that were oppositely directed with regard to gaze direction. Moreover, amplitudes were decreased in vertical EOG channels bilaterally with stimulation along the naso-occipital (x) axis in supine and prone positions.

Healthy subjects had a normal VCR bilaterally, whereas it was attenuated or absent ipsilaterally in all patients with UVD and bilaterally in all patients with BVD.

DISCUSSION

Vestibular response amplitudes (up to 100%), elicited by vibrations can have even in normal subjects, considerable inter-side differences and the amplitude difference can approach 100%.20 This may be due to CFA, which can be invisible to the naked eye, or to hidden remnants of past vestibular insult. Thus the findings of the current study with regard to the control group as well as the second heterogeneous UVD group with unspecified vestibular pathology might be biased due to the selection criteria based on the findings of the VCR elicited by head taps. However, the findings in the first UVD group with anatomically proved complete vestibular deafferentation were undoubtedly consequences of the vestibular disorder.

Passive oscillatory responses and lid artefacts

To determine the origin of the recorded events the passive movements of electrodes, eyelids, and the globe should be taken into account. The natural frequency of oscillation for orbital tissues is above 12 Hz with a resonance frequency in the range of 50–63 Hz, so vibrations applied to the skull could induce passive eye and lid movements.33 The amplitudes of the responses were in the range of values measured in vibration induced passive eye movements.34 Tonic innervation and ageing may change the viscoelastic properties of the eye muscles and modulate the resonance frequency and amplitude. Thus the absence or diminution of the responses in cadavers as well as individuals with palsies of the extraocular muscles and BVD could not exclude the possibility of passive oscillatory origin of responses. Indeed all but one of the currently presented patients with BVD were of advanced age.

Vertical EOG records always reflect an interaction of the eye and lid movement. The eyelid acts as sliding electrode, increasingly shunting the positive corneal pole to the upper EOG electrode while covering the larger surface of the cornea, or moving downward.36 In all vertical eye movements, the lids follow the globes closely but small differences between lid
Brainstem reflexes

Blink reflex

Only the trigeminofacial blink reflex shows a short latency R1 component possessing similar latency to the EOG responses in this study. This early component can generally be elicited only by stimulating the first division of the trigeminal nerve, which was avoided in this study.\(^{46}\) On the other hand, the later occurring symmetrical R2 component can be elicited by various sensory stimuli including the stimulation of sensory branches beyond the trigeminal territory. The latency of this component generally exceeds 20 ms.\(^{47,48}\) The R2 component is invariably accompanied by inhibition of levator muscle activity, as well as disconjugate medially and downward directed eye movement, but the orbicularis oculi response always precedes the associated eye and lid movements.\(^{37,50}\) Since in the present study, the responses in vertical EOG were always of higher amplitude on the side of lesioned labyrinth, even in the stage of complete facial palsy after surgery, and were absent or substantially diminished in patients of profound BVD, the responses recorded were not in any way generated by voluntary or reflex blinking.

Tonic vibration reflex of the jaw closure muscles

Primary Ia spindle afferents are extremely sensitive to vibration, and thereby tonic vibration reflexes could have been elicited in certain cranial muscles in this study. Although facial muscles lack muscle spindles, the jaw closing muscles are richly endowed. The physiology of the masseter muscle tonic vibration reflex has a unique electromyographic pattern of synchronised waves with a one to one relationship with the vibration cycles.\(^{46}\) EOG electrodes might naturally pick up the vibration not only elicits a tonic vibration reflex but also potentiates the phasic reflexes. Moreover, the jaw closure tonic vibration reflex has some peculiar features due to lack of presynaptic inhibition onto spinal afferents. Unlike in limb muscles, vibration not only elicits a tonic vibration reflex but also potentiates the phasic reflexes. Nevertheless, the tonic vibration reflex evolves gradually with a rather long latency, because it depends on progressive facilitation and recruitment in polysynaptic proprioceptive pathways.\(^{46}\) Therefore it is unlikely to appear in the first several tens of milliseconds following the single vibrational impulses applied at a low repetition rate of one to three per second, regardless of the stimulus intensity.

Proprioceptive reflexes of the extraocular muscles

The ability of the oculomotor system to determine eye position is essential. According to earlier data the control of eye movement and position appeared to be primarily effently coded. Indeed, there is no swift stretch reflex for eye muscles despite a generous complement of muscle spindles.\(^{34}\) Nevertheless, there is abundant evidence that the brain uses information from eye muscle proprioceptors. The palisade endings associated with the tips of the multiply innervated non-twitch muscle fibres are the most likely receptors in the principal sensory apparatus of the extraocular muscles. They are innervated by tonic motor neurones with small diameter axons, which mediate signals related only to intended eye positions. Thus they may participate in a proprioceptive system important for setting and stabilising the alignment of the eye.\(^{46}\) However, due to slower conduction and execution
time the motor responses mediated by this system could be expected to occur at longer latencies as the responses in the current study.

Cervico-ocular reflex

Skull vibrations are also conducted to cervical muscles and activate their la afferents, thereby eliciting the cervico-ocular reflex. Electrophysiological experiments suggest that this reflex is mediated via the vestibular nuclei but the precise projection is only partially known.27 The cervico-ocular reflex is enhanced in patients with vestibular areflexia because of the increased central weighting of somatosensory neck information, which substitutes for missing vestibular input.48 49 It is likely that pathways that mediate this reflex are polysynaptic, because this reflex does not occur with latency shorter than 40 ms.49 Nevertheless, if the short latency events recorded in the current study were under the control of vestibular nuclei, the possibility of their conditioning by cervicovestibular input could not be rejected definitively.

Vestibulo-ocular reflexes

Skull as well as neck vibrations can evoke ocular movements and it is controversial whether these eye movements are caused by activation of vestibular or cervico-ocular reflexes.47–49 Since VCR with disynaptic latency can be activated by bone conducted vibrational impulses upon the skull, it is unlikely for the same stimuli not to simultaneously also elicit disynaptic VOR. Close behavioural coupling of the VCR and VOR is necessary and mammals without such coupling could not withstand natural selection pressure.

Similar eye movements to the responses recorded in the current study have been obtained by galvanic stimulation of the whole vestibular nerve, by electrical stimulation of the utricular nerve, the ampullary nerves of the semicircular canals or vestibular end organs, and by acoustic clicks as well as natural vestibular stimulation.50–56

As a consequence of low frequency vibrations the head moves as a whole, executing parallel, or so-called translational movements to and from the site of application of vibration. Hence, the vestibular end organs embedded in the temporal bone are alternately subjected to inertial pressure.48 If the vibration pulse is realised as a sine wave, then the dominant component of the recorded events could be considered a response to a second phase of this wave probably due to the asymmetry in the otolith transfer functions.57 58

An important finding of the current study is the striking asymmetry in bidirectional sensitivity and this non-linearity is even more marked at the level of second order neurones.59 However, this asymmetry is more complex for the otolith system since it prefers oppositely directed hair cell deflections during dynamic translations versus static tilts and off-centre rotations.60 61 Thus the enhancement of the response in vertical EOG channel on the side of vestibular insult and/or diminution on the healthy side may be a consequence of the directional asymmetry of the remaining unbalanced labyrinthine input.

Latencies of responses about 3–15 ms, with low inter-individual variability in this study are in accordance with those found for the disynaptic aVOR.62–64 We have not made much effort to demonstrate the distance effect, because it was felt the technique was not suitable for detection of this kind of change. Even with more natural stimuli, due to the exponential shape of VOR response, the robust distance effect was very subtle at the onset. The rate of the gain adjustment for viewing distances of 40 cm versus 10 cm at the onset was found to differ only by one standard deviation in velocity traces using the search coil technique.6 Hence, it was easily lost in the raw EOG eye position records. On the other hand, the responses were much higher than those required for an ideal compensatory VOR and the lid artefacts could have contributed to them. Nevertheless, for high frequency stimuli used in this study, the gain of the response was primarily determined by the frequency of the stimuli, whereas some kinematical requirements of the response may have been attenuated. Apart from the vertical gaze effects on the responses that were caused by lid artefacts, the other kinematical aspects and positional effects might be consistent with otolith driven reflexes.65 The increase in amplitude of the response in vertical EOG channel on the side of adducting eye, and the decrease on the side of abducting eye, could be compatible with subsequent activation of both oblique muscles, yet evidence for disynaptic connections from otolith organs to contralateral inferior oblique muscle is lacking.66–68

Finally, the apparently disconjugate responses may even reflect the basic monocular organisation of the oculomotor system. Indeed, during sleep, rapid eye movements are found to be monocular or disjunctive, and electrophysiological studies in primates have revealed that pre-motor position–vestibular-pause neurones fire in relation to monocular eye position, rather than to conjugate eye movements.69–71

CONCLUSIONS

In the present study, EOG responses to skull vibration were generated either by phasic VOR or by passive oscillatory movements of the orbital tissues. Even if the latter were true, there was a striking asymmetry in the responses of vertical EOG channels elicited by stimuli along the naso-occipital (x) axis in patients with UVD compared with normal subjects. This finding was at least in accordance with the movement of the eyes on the healthy and lesioned side from different starting positions, due to otolith mediated sustained skew torsion in patients with UVD. This asymmetry provides a consistent, longlasting lateralising clue for testing at least complete UVD. We are aware of several drawbacks of our study due to the limited reliability of the methods of recording and analysis. However, despite these limitations the findings provide at least circumstantial evidence for the possibility of investigating VOR using vibrational impulse stimuli.

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A case of late onset sporadic Parkinson’s disease with an A53T mutation in α-synuclein

Parkinson’s disease is a common progressive neurological disorder characterised by loss of nigral dopaminergic neurones. Rare autosomal dominant familial cases have been associated with point mutations in α-synuclein, but the vast majority of cases occur sporadically in older patients without an obvious cause. We now report a unique case of typical late onset Parkinson’s disease without a family history which was associated with an A53T mutation in α-synuclein.

Case report

A war veteran of Polish origin was initially referred for assessment of his parkinsonian condition in 1997 at the age of 74. At presentation his history was of progressive bradykinesia, difficulty in rising from his chair, a tendency to fall, and mild tremor. He was a smoker and had been treated for hypertension and hypercholesterolaemia, but he gave no clear history of cerebrovascular disease. His Austrian mother died at 91 years of age and his French father lived to 89, neither suffering from symptoms of Parkinson’s disease. He had four brothers and three sisters, none of whom had symptoms of Parkinson’s disease (two died in their 20s during the war, the others died at ages 68, 76, 78, and 86, and one has lost touch with the family). Furthermore, his three children, each now in their sixth decade, currently have no diagnosis of Parkinson’s disease. The family know of no relatives of Italian or Greek origin.

Examination was consistent with Parkinson’s disease, with a typical shuffling gait, bilateral cogwheel rigidity, and mild tremor, but no pyramidal or cerebellar signs. Investigations were normal, but magnetic resonance imaging of his brain was not possible because of metal shrapnel in his left orbit, face, and nose from the second world war. Computed tomography of the brain showed only mild age related cerebral atrophy without evidence of vascular disease.

Before presentation he had been prescribed co-beneldopa 62.5 mg three times a day with some symptomatic benefit. In July 1997 a five week trial off levodopa caused a deterioration in his symptoms, therefore his co-beneldopa was restarted and increased to 125 mg three times daily, and selegiline was started. On this treatment his symptoms remained stable for the next three years, and a second trial without levodopa or a dopamine agonist was attempted in May 2000, which again provoked marked bradykinesia and deterioration in his gait. His treatment was restarted after only five days, following which the symptoms once again resolved, showing the clear levodopa responsive element to his condition. He died in August 2002 but a necropsy examination of the brain was not undertaken.

Genetic analysis

PCR primers were designed from 5’ untranslated region (UTR) and 3’UTR spanning each exon of α-synuclein (from NACP/α-synuclein sequences submitted to NCBI database; accession number U46896-U46901; primer pairs designed to amplify exon 3 were 3F: GAGACCTCTGTTAGCTGG, and 3R: GACTGATATGTTCTTAGATGCTG. Polymerase chain reaction (PCR) products were purified using QIAquick columns and sequenced according to the manufacturer’s protocols by dye terminator (BigDye) methods using an ABI 377 automated sequencer (Applied Biosystems, Foster City, California, USA). All sequences were edited and confirmed by entering them into the BLAST algorithm database at the National Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov/blast).

A single mutation from G to A at nucleotide 209 in exon 3 of the α-synuclein gene from this patient (fig 1A). The sample was reanalysed twice following this observation, and also verified in the reverse direction. Restriction digest of the PCR products was carried out with Tsp45 I (New England BioLabs, Beverly, Massachusetts, USA), and digested products were separated by electrophoresis on a 2% agarose gel (fig 1B). The results correspond to an alanine to threonine shift at position 53 of α-synuclein (an A53T mutation).

Comment

The discovery of families with autosomal dominant Parkinson’s disease together with the subsequent development of symptomatic α-synuclein transgenic models of the disease has provided strong support that a point mutation in α-synuclein is sufficient to cause this disorder. Furthermore, the discovery of α-synuclein in the Lewy bodies of patients with the common sporadic form of Parkinson’s disease has suggested that this protein may well play a central role in all forms of the disease.

Until now the A53T mutation in α-synuclein has only been described in a large kindred of Italian origin and a small number of unrelated families of Greek origin, each with autosomal dominant inheritance of the Parkinson’s disease phenotype. The clinical phenotype and response to levodopa in the Italian kindred is relatively typical for Parkinson’s disease, but with an earlier mean age of onset at 46 (SD 13) years, and a relatively rapid course, averaging 9.7 years from onset to death.

The case we describe had no such family history, first noticed symptoms in his eighth decade, and had a clinical phenotype compatible with sporadic idiopathic Parkinson’s disease. He had a relatively mild tremor, as noted in the Italian kindred. This case is therefore of particular interest as it appears to represent a unique sporadic mutation in α-synuclein that was not found in the remaining 60 control and patient samples that we analysed, and has not been reported in other series. Furthermore, despite carrying the
genetic mutation this patient developed symptoms much later in life than most members of the Italian kindred described above.

The penetrance of the A53T α-synuclein gene in the Italian kindred has been estimated at 85% so, although unlikely, it is theoretically possible that in this case the mutant gene is asymptomatic and the patient has developed unrelated late onset sporadic Parkinson’s disease. Alternatively it is possible that the relatively mild late onset of Parkinson’s disease in this patient represents a dose effect of the mutant gene. For example, both duplication and triplication of the α-synuclein gene locus have recently been found to cause familial Parkinson’s disease, and the different severity of clinical phenotype seems to be correlated with the dose of α-synuclein.

This unique case extends the repertoire of patients in whom Parkinson’s disease is associated with point mutations of α-synuclein. While we do not advocate routine clinical screening, this case suggests that further evaluation of mutations on this gene should be considered in cases of sporadic Parkinson’s disease.

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References


Decreased CSF hypocretin-1 (orexin-A) after acute haemorrhagic brain injury

The novel hypothalamic neuropeptides orexins, or hypocretins, have gained much attention as potential modulators of various different physiological functions. Deficient orexin neurotransmission may be responsible for excessive somnolence, as shown in several conditions related to secondary narcolepsy, through direct or indirect damage to the posterior hypothalamus and its connections. We have studied for the first time the longitudinal changes of hypocretin-1 concentrations in cerebrospinal fluid (CSF) in patients with acute haemorrhagic brain injury.

Nine patients from a previously reported cohort and 21 controls (seven women and five men, median age 38 years, range 17 to 70) with other neurological disease were enrolled in the study (table 1). The patient group included five subjects with intracerebral haemorrhage and five with subarachnoid haemorrhage. All patients had extraventricular drains inserted within a median of two days of disease onset (range 2 to 36) as a treatment procedure because of increasing signs of hydrocephalus. Morning CSF samples were collected twice: first between day 1 and day 2 after catheter insertion, and second between day 4 and day 10. Patients were assessed using the Glasgow coma score (GCS) at presentation and the Glasgow outcome score (GOS) after three months.

The control group consisted of two patients with primary dementia, two with chronic headache, one with ataxia syndrome, and seven with non-specific neurological symptoms. None reported sleep abnormalities. All samples were stored at ~80°C until analysis. Hypocretin-1 was measured blind to diagnosis by a direct radioimmunoassay of 100 μL of CSF (Phoenix Pharmaceuticals, Belmont, California, USA; detection limit 40 pg/ml, intra-assay variation <3%), as described previously. Samples with undetectable concentrations (value below 40 pg/ml) were operationally plotted at 0 pg/ml. Statistical tests were carried out with the GraphPad InStat 3.05 software package using the non-parametric Mann–Whitney U test.

There was a significant difference in median CSF hypocretin-1 concentrations between the controls (319.4 (302 to 361) pg/ml) and acute brain injury patients (100.4 (0 to 145.2) pg/ml) (calculated from the means of two measurements; median (range)). No difference was found for sex or age (table 1). The concentration of hypocretin-1 in CSF on the first day of sample collection (24 hours after catheter insertion) was 98.8 (0 to 147) pg/ml and did not change significantly over the observation period (114.0 (0 to 144) pg/ml). All concentrations were either lower than control values (>200 pg/ml), in the intermediate range (110 to 200 pg/ml), or in the very low, narcolepsy range (<110 pg/ml). Two patients (Nos 4 and 6) had undetectable levels, while all others had moderately decreased values compared with the cut off level of 196 pg/ml. In patient 6, the hypocretin-1 level increased to 53 pg/ml, but it remained undetectable in patient 4. Both patients suffered from spontaneous intracerebral haemorrhage, which was localised either to the thalamus (patient 4) or to the midbrain (patient 6). Other patients with moderately decreased hypocretin-1 levels were diagnosed as having subarachnoid haemorrhage (six patients) and intracerebral haemorrhage in the frontal lobe (one patient). There was no correlation between hypocretin-1 levels and GCS at disease onset or GOS at three months after disease onset.

Comment

This is the first study to show decreased levels of hypocretin-1 in the CSF of patients in the first week after acute brain injury caused by haemorrhagic stroke. Our data are in line with previous observations in patients with traumatic brain injury.

The findings seem important in the light of a study investigating long term outcome in patients after subarachnoid haemorrhage. More than 75% of patients reported excessive fatigue or daytime sleepiness, which persisted for long period (months to years) after the event. However, the exact mechanism responsible for the complaints remained unknown. Although the aetiology may be quite complex, an abnormality in the hypocretin/orexin system could make an important contribution to the phenomenon.

How intracerebral haemorrhage affects hypothalamic function remains obscure. In two of our patients direct damage to the thalamus/brain stem circuits appeared to be responsible. However, the remainder of the patients had a subarachnoid bleed or lesions in remote brain structures unrelated to the

Table 1 Patient characteristics

<table>
<thead>
<tr>
<th>Case</th>
<th>Sex</th>
<th>Age (y)</th>
<th>Diagnosis</th>
<th>Lesion and complications</th>
<th>GCS</th>
<th>IVS</th>
<th>GOS</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>M</td>
<td>45</td>
<td>ICH</td>
<td>HC</td>
<td>6/15</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>2</td>
<td>M</td>
<td>66</td>
<td>SAH (ACA aneurysm)</td>
<td>HC, multiple infarcts</td>
<td>5/15</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>3</td>
<td>M</td>
<td>54</td>
<td>SAH (PCA aneurysm)</td>
<td>HC, occipital infarct, re-bleed</td>
<td>3/15</td>
<td>15</td>
<td>15</td>
</tr>
<tr>
<td>4</td>
<td>F</td>
<td>59</td>
<td>ICH (spontaneous)</td>
<td>HC, thalamus</td>
<td>7/15</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>5</td>
<td>M</td>
<td>59</td>
<td>SAH</td>
<td>HC</td>
<td>14/15</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>6</td>
<td>F</td>
<td>59</td>
<td>ICH (spontaneous)</td>
<td>HC, midbrain, re-bleed</td>
<td>6/15</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>7</td>
<td>M</td>
<td>72</td>
<td>SAH</td>
<td>HC</td>
<td>14/15</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>8</td>
<td>F</td>
<td>70</td>
<td>SAH (ACA aneurysm)</td>
<td>HC, re-bleed</td>
<td>15/15</td>
<td>8</td>
<td>5</td>
</tr>
<tr>
<td>9</td>
<td>M</td>
<td>45</td>
<td>ICH (spontaneous)</td>
<td>Frontal lobe</td>
<td>8/15</td>
<td>36</td>
<td>36</td>
</tr>
</tbody>
</table>

ACA, anterior cerebral artery; F, female; GCS, Glasgow coma scale (on admission); GOS, Glasgow outcome scale (after 3 months); HC, hydrocephalus; ICH, intracerebral haemorrhage; IVS, intraventricular shunt (days after disease onset); M, male; PCA, posterior cerebral artery; SAH, subarachnoid haemorrhage; y, years.
hypothalamus. Previously we provided evidence of diffuse axonal injury in these patients and speculated that this represented one mechanism of disruption of the hypothalamic circuits. A remote chemical mechanism contributing to the decrease in hypocretin-1 levels in the CSF. It is well recognised that a large amount of blood entering the CSF compartment or brain parenchyma produces neurotoxic effects through various different mechanisms, including oxidative haem and iron metabolism and secondary oedema with abnormalities of brain perfusion.

The decreased hypocretin-1 concentrations in our patients might have resulted from the dilution effect of the bleed into the CSF with the development of secondary hydrocephalus in the course of their disease. However, this seems rather unlikely as intraventricular drainage was initiated early and the first samples were collected at least 24 hours after catheter insertion. One might also expect that in hydrocephalus complicating subarachnoid haemorrhage there would be an accumulation of CSF constituents owing to reduced absorption by subarachnoid villi, which would go against our hypothesis; however, an opposite effect was observed and this persisted during the course of the disease.

An important caveat of our study is that control samples were obtained by lumbar puncture, and lumbar CSF is likely to have a different composition from cisternal CSF. However, the concentrations of different neurotransmitters in the ventricular CSF are reported to be higher than in corresponding lumbar puncture specimens.

Further studies are needed to investigate prospectively the relation between hypocretin-1 production and sleep–wake cycle abnormalities in patients after haemorrhagic stroke.

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**REFERENCES**


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**BOOK REVIEW**

The essential handbook of memory disorders for clinicians


Any reader who is familiar with the editors’ comprehensive and authoritative “Handbook of Memory Disorders” will experience a powerful sense of déjà vu on opening this volume, and may be forgiven for wondering what exactly the point of it is. The Readers’ Digest edition? Neuropsychology lite? With its 392 pages this is hardly a pocket companion. The real reason can be found in an (extremely brief) preface: the 35 chapters of the original work had resulted in a “heftier and more expensive book, which might well be seen as less directly relevant to clinical practice”. In other words, a self-confessed case of “mega biblion mega kakon”, and doubtless a publisher’s marketing wheeze.

A wizard one? Perhaps. Thirty-five chapters have been whittled down to 15, of which all but one have direct clinical relevance, ranging from the amnesias of childhood to a review of rehabilitative strategies for the memory impaired. The exception is Baddeley’s opening essay on contemporary and historical views of the psychology of memory. While such an overview is by no means out of place, its theoretical emphasis might perhaps have been acknowledged by according it the title and status of Introduction rather than merely “Chapter 1”.

The remaining contents are also rather arbitrarily ordered, and the clinician in search of an up-to-the-minute review of some aspect of diagnosis or management is hardly guided by it to the most relevant pages. All the important themes—evaluation, differential diagnosis, management—are well represented, but needlessly interleaved. Chapters dealing with the assessment of memory disorders and the distinction between disorders of memory and other cognitive systems come after those in which specific subtypes of memory dysfunction are discussed. No fewer than four chapters discuss remediation and rehabilitation, while the discussion of retrograde amnesia is entirely subsumed within a review of psychogenic disorders, and the critically important topic of semantic memory is completely neglected.

So while this handbook will undoubtedly be of interest to clinicians, I suspect that many will prefer to distil the essence of the subject from the more comprehensive parent volume, and regard the additional heft and cost as a price worth paying.

P Garrard

**CORRECTION**

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Jombik P and Bahyl V. Short latency responses in the averaged electro-oculogram elicited by vibrational impulse stimuli applied to the skull: could they reflect vestibulo-ocular reflex function? *J Neurol Neurosurg Psychiatry* 2005;76:222–8). The first sentence of figure 1 legend should read: Averaged electro-oculography (EOG) responses in a normal subject elicited by stimuli along the interaural axis.