

restored and he found himself lying on the floor. On recovery there was no confusion, drowsiness, dysphasia, or diuresis. Often, however, he sustained soft tissue injuries to his face and scalp.

Witnesses reported that the patient would, without warning, suddenly collapse to the ground where he would remain unrousable, inaccessible, and motionless for 90 to 120 seconds. On two occasions he appeared confused and disorientated immediately before a collapse. During the period of unconsciousness he would demonstrate no involuntary movements, orofacial automatisms, or cyanosis but he would become pale and "ashen" while staring straight ahead with a glazed look. On resolution of the episode his colour would return to normal and within 2 minutes he would have fully recovered. Unusually during one reported episode of unconsciousness he was seen to briefly extend the fingers of both hands.

He was admitted to his local hospital and CT, MRI, interictal EEG, and 24 hour ECG were normal. No episodes were witnessed while he was an inpatient but they were thought to be epileptic in origin and therefore he was started on phenytoin, with no benefit. Carbamazepine was added, again with minimal effect.

The patient was then referred to the Epilepsy Assessment Centre of The National Society for Epilepsy and National Hospital for Neurology and Neurosurgery for further investigation and management.

Cardiovascular and neurological examination was normal, as were MRI and routine interictal EEG. Sixteen channel ambulatory EEG using an Oxford Instruments digital EEG receiver was performed continuously for 340 hours before an episode was captured. Interictally rare spikes were seen over the right frontocentrottemporal region during sleep. The onset of the episode was not witnessed and the patient was found lying on the floor, regaining consciousness at about 07:06. The event EEG showed a short run of bilateral semirhythmic 2-3 Hz activity at 07:04:34 (figure A), persisting for 8 seconds before being obscured by muscle and movement artefact. Twenty four seconds after the first EEG change, at 07:04:58, the ECG changed from sinus rhythm at 90 bpm to a brief period of sinus bradycardia, followed by a period of asystole with only very occasional ventricular complexes lasting 25-30 seconds (figure B). After a few seconds of bradycardia then tachycardia, sinus rhythm was restored. Throughout the episode the QT interval on the ECG remained within normal limits. The EEG became visible again 16 seconds into the asystolic period, at which time it was dominated by diffuse low amplitude slow activity at <1-2 Hz which persisted for 10 seconds (figure C). This was followed by marked attenuation of the EEG activity over the next 10 seconds before large amplitude generalised rhythmic <1Hz activity became apparent. Diffuse theta activity was seen for a further 15 seconds before the EEG returned to its resting state.

A VVI permanent pacemaker was inserted. The phenytoin was withdrawn and replaced by lamotrigine. Carbamazepine was left unchanged. The patient was discharged, his medication left unaltered, and at follow up 9 months later reported no further episodes.

Cardiac dysrhythmias are an uncommon but serious consequence of partial seizures. Our case is unusual because of the duration of the asystole. In a series of 26 patients with 74 temporal lobe seizures in which simulta-

neous EEG and ECG recordings were acquired, ictal arrhythmias occurred in 52% of seizures, the commonest being irregular abrupt changes in heart rate, (both acceleration and deceleration) occurring towards the end of the period of EEG abnormality.¹ Interictally, patients with epilepsy seem no more likely than age and sex matched healthy subjects to experience arrhythmias although in one study patients with epilepsy had a faster ventricular rate and a longer QT interval than control subjects.³

It has been hypothesised that there is lateralisation with respect to central autonomic cardiac control with an increase in heart rate seen after an intracarotid injection of amobarbital and inactivation of the left hemisphere and a decrease in heart rate on right hemispheric inactivation. Experimental stimulation of the rostral posterior insular cortex in anaesthetised rats has been shown to induce tachycardia and more caudal region stimulation to cause bradycardia.⁴ Additionally, prolonged stimulation resulted in ventricular ectopics, heart block, QT prolongation, and death. In presurgical temporal lobectomy patients stimulation of the left insular cortex (particularly posteriorly) produced bradycardia and a depressor response significantly more often than tachycardia and a pressor effect.⁵ It was suggested that an epileptic discharge in the insular cortex may result in cardiac arrhythmias.

Recurrent episodes of loss of consciousness are a common clinical problem. An accurate diagnosis relies principally on the patient's and witnesses' accounts of events. Further investigations are frequently required which are often normal unless an episode is captured during monitoring. Recording solely the EEG or the ECG may result in erroneous conclusions being drawn and insufficient or inappropriate therapy being instituted. Distinction between a primary cardiac arrhythmia and a secondary central arrhythmia is possible only with simultaneous EEG/ECG recordings.

FERGUS J RUGG-GUNN
JOHN S DUNCAN
SHELAGH J M SMITH

Epilepsy Research Group, University Department of
Clinical Neurology, Institute of Neurology, The
National Hospital for Neurology and Neurosurgery,
Queen Square, London WC1N 3BG, UK

Correspondence to: Professor John S Duncan,
National Society for Epilepsy, Chalfont St Peter,
Gerrards Cross, Bucks SL9 0RJ, UK
email j.duncan@ion.ucl.ac.uk

- 1 Blumhardt LD, Smith PEM, Owen L. Electrocardiographic accompaniments of temporal lobe epileptic seizures. *Lancet* 1986;i:1052-5.
- 2 Reeves AL, Nolle KE, Klass DW, et al. The ictal bradycardia syndrome. *Epilepsia* 1996;37:983-7.
- 3 Drake M, Reider C, Kay A. Electrocardiography in epileptic patients without cardiac symptoms. *Seizure* 1993;2:63-5.
- 4 Oppenheimer SM, Hachinski VC, Cechetto DF. Cardiac chronotropic organization of the rat insular cortex. *Society of Neurosciences Abstracts* 1989;15:595.
- 5 Oppenheimer S, Gelb A, Girvin J, et al. Cardiovascular effect of human insular cortex stimulation. *Neurology* 1992;42:1727-32.

Respiratory insufficiency in a patient with hereditary neuropathy with liability to pressure palsy

Hereditary neuropathy with liability to pressure palsies (HNPP) is an autosomal dominant disorder, the molecular basis of which is a 1.5 mb deletion in chromosome 17p11.2 including the peripheral myelin protein-22

(PMP-22) gene.¹ HNPP typically presents recurrent pressure palsies of peripheral nerves, such as the axillary, median, radial, ulnar, or peroneal nerves, at common entrapment sites. Respiratory muscle weakness has not been previously reported in HNPP. We describe a patient with HNPP in whom respiratory failure and proximal muscle weakness were prominent features.

The patient started to have dyspnoea on exertion at the age of 44. At the age of 47, he noticed a slowly progressive weakness of the pelvic girdle and lower limbs. At the age of 57, he experienced difficulty in going up stairs. However, he was almost independent in daily life. At the age of 60, he was admitted to Narita Red Cross Hospital as an emergency patient with a coma due to CO₂ narcosis (PCO₂ 117.6, PO₂ 64.0). Responding to mechanical ventilatory support, he completely recovered consciousness within a day. His respiratory condition in the daytime improved to that previously. However, he needed mechanical ventilation during sleep because of nocturnal hypoventilation.

The patient had no history of diabetes mellitus, pulmonary disease, or other medical problems. There was no familial history of neurological disorder, including entrapment neuropathies. After a few months, he noted that in his teens he had experienced some episodes of right peroneal and right axillary nerve palsies which resolved themselves over a few months.

In a neurological examination, the patient's mental state and cranial nerves were normal. Evidence of muscular atrophy and lumbar lordosis was found. The muscular atrophy was prominent in the shoulder girdle, intercostal muscles, paravertebral muscles, and pelvic girdle, and moderate atrophy was present in all four limbs (figure). There was moderate weakness of the shoulder and pelvic girdle and mild weakness of the distal limbs. The thorax showed poor respiratory movement, and the patient showed paradoxical movement of the abdomen in the supine position. Tendon reflexes were hypoactive in all limbs. The patient's sensations of touch and pain were mildly impaired in the four distal limbs. His position sensation was normal. His vital capacity was 1.9 l (55% of the normal mean) in the sitting position, but 1.3 l (38%) in the supine position. The percentage of forced expiratory volume in 1 second was normal (99%). Chest radiography at inspiration and expiration showed poor movement of the diaphragm but no abnormality in the lung field. Routine haematological and serological studies gave normal results. No monoclonal or polyclonal proteins were detected. IgG and IgM antibodies to gangliosides GM1 and GD1b were negative. Analysis of CSF showed 1 lymphocyte/mm³ and 25 mg/dl protein. Motor nerve conduction studies showed prolonged distal latencies in the right median (8.8 ms (normal value in our laboratory <4.6)) and ulnar (6.2 ms (normal <3.6)) nerves, and moderate decreased conduction velocities in the right median (40 m/s (normal >45)), ulnar (45 m/s (normal >49)), tibial (35 m/s (normal >38)), and peroneal (29 m/s (normal >41)) nerves. There were moderate decreases in the amplitude of compound action potentials in all the nerves tested, and an amplitude reduction of 50% was detected across the cubital tunnel of the right ulnar nerve. Minimum F wave latencies were prolonged in all the nerves tested. The latency in the right phrenic nerve was slightly



General muscle atrophies, which are most prominent in the trunk are shown. A tracheotomy was performed for nocturnal hypoventilation because the patient required mechanical respiratory support during the night.

delayed (8.7 ms (normal<8.0)). Sensory nerve conduction studies showed a reduced amplitude of sensory nerve action potentials and conduction slowing in all the nerves tested. Electromyography carried out in the supraspinatus, deltoid, biceps, flexor carpi ulnaris, brachioradialis, quadriceps femoris, biceps femoris, tibialis anterior, and gastrocnemius muscles showed polyphasic motor unit potentials of long duration, but denervation potentials were rare. A left sural nerve biopsy showed scattered tomaculous thickening of the myelin sheath and some abnormally thin axonal myelin sheaths. The density of myelinated fibres was reduced (5726/mm²). A gene analysis disclosed a 53% gene dose of PMP-22 related to normal controls, using Southern blots of DNA digested with EcoRI.

Given the possibility of superimposing demyelinating neuropathy, especially chronic inflammatory demyelinating polyneuropathy, oral prednisolone (60 mg/day) was given for 1 month. However, the patient's clinical condition did not respond to this treatment. Pulmonary dysfunction and proximal muscle weakness were almost steady during the next 3 years.

We examined the patient's elder sister (64 years old), elder brother (62 years old), and younger sister (58 years old), although they had no neurological complaints. All of them had experienced generalised hyporeflexia or areflexia but no weakness or sensory loss, and nerve conduction studies showed moderate conduction slowing with accentuation at the common entrapment sites, suggesting demyelinating neuropathy.

Our patient recalled experiencing recurrent episodes of transit entrapment mononeuropathies, and the familial occurrence of asymptomatic entrapment neuropathy was detected by nerve conduction studies. The

presence of tomacula, and genetic analysis confirmed a diagnosis of HNPP. However, the patient's dominant clinical features—respiratory failure and proximal muscle weakness—were atypical for HNPP. Although respiratory muscle weakness has been reported in hereditary motor and sensory neuropathy (HMSN),²⁻⁴ there has been no report of respiratory insufficiency associated with HNPP to our knowledge.

The weakness of the truncal muscles, including the respiratory accessory muscle, is a possible cause of respiratory failure in our patient. On the other hand, he had experienced hypoventilation in the supine posture and paradoxical movement of the abdomen, which suggested diaphragmatic weakness.² Also, chest radiography showed poor movement of the diaphragm. Although the prolongation of distal latency in the phrenic nerve was mild considering the severity of respiratory failure, assessment of axonal loss is not possible with phrenic nerve stimulation. In fact, phrenic nerve latency is not necessarily associated with pulmonary dysfunction in HMSN.⁴

Diffuse proximal weakness in our patient is an uncommon finding as for HNPP. Mancardi *et al*⁵ reported on three patients with progressive sensory-motor polyneuropathy associated with 17p11.2 deletion, and the initial symptom of one patient was proximal weakness in one arm. We propose that our patient represents a clinical phenotypic variability among HNPP. It may be necessary to pay attention to respiratory function in HNPP.

We thank Dr T Yamamoto from the University of Occupational and Environmental Health for the gene analysis and Mr T Nagase from Chiba University for his technical help with the sural nerve biopsy.

MASATO ASAHINA
SATOSHI KUWABARA
TAKAMICHI HATTORI
Department of Neurology, School of Medicine,
Chiba University, 1-8-1 Inohana, Chuo-ku,
Chiba 260-8670, Japan

MASATO ASAHINA
KAORU KATAYAMA
Department of Neurology, Narita Red Cross Hospital,
90-1 Iida-cho, Narita-shi, Chiba 286-0041, Japan

Correspondence to: Dr Masato Asahina, 1-8-1 Inohana, Chuo-ku, Chiba-shi, Chiba 260-8670, Japan. Telephone 0081 43 222 7171 ext 5414; fax 0081 43 226 2160; email: masatoasahina@msn.com

- 1 Chance PF, Alderson MK, Lepping KA, *et al*. DNA deletion associated with hereditary neuropathy with liability to pressure palsies. *Cell* 1993;72:143-51
- 2 Eichacker PQ, Spiro A, Sherman M, *et al*. Respiratory muscle dysfunction in hereditary motor sensory neuropathy, type I. *Arch Intern Med* 1988;148:1739-40.
- 3 Hardie R, Harding AE, Hirsch N, *et al*. Diaphragmatic weakness in hereditary motor and sensory neuropathy. *J Neurol Neurosurg Psychiatry* 1990;53:348-50.
- 4 Carter GT, Kilmer DD, Bonekat HW, *et al*. Evaluation of phrenic nerve and pulmonary function in hereditary motor and sensory neuropathy. *Muscle Nerve* 1992;15:459-62.
- 5 Mancardi GL, Mandich P, Nassani S, *et al*. Progressive sensory-motor polyneuropathy with tomaculous changes is associated to 17p11.2 deletion. *J Neurol Sci* 1995;131:30-4.

Spinal accessory neuropathy and internal jugular thrombosis after carotid endarterectomy

Spinal accessory neuropathy is a rare complication of carotid endarterectomy (CEA).¹ Internal jugular venous thrombosis after CEA has also been reported rarely, but is likely more common; as internal jugular