“Sporadic” familial amyloidotic polyneuropathy in a German patient with B cell lymphocytic leukaemia

We report a 70 year old German man presenting with a three year history of progressive numbness and painful tingling and burning paraesthesiae in his hands, feet, and lower legs, which had worsened during recent months. He also reported unsteadiness of gait, fatigue, night sweats, loss of appetite, and a weight loss of 12 kg within one year. He denied bowel or bladder problems and alcohol ingestion, but admitted smoking (110 pack-years). His family history was negative for neurological diseases.

The family was originally from Gdansk (now Northern Poland). The patient’s father and his four siblings all reached their 80s without developing neurological symptoms. The patient’s mother died at the age of 64 of blood cancer, and her half brother died at the age of 78. Three of the patient’s siblings died at the ages of 1, 17, and 33 (starvation, killed in the war, stomach cancer). Two further sisters, aged 64 and 69, their descendants, and the patient’s own five sons and their children were healthy. Both of the patient’s grandmothers died in their 80s, whereas the paternal grandfather died early of unknown cause, and the maternal grandfather drowned in his 30s.

Neurological examination revealed severe ataxia of gait and stance, atrophy of the small hand and foot muscles, and bilateral distal pareses (3–4/5 on the MRC scale), diminished tendon jerks, a glove and stocking distribution of hypoaesthesia for all sensory qualities up to the mid-thighs and elbows, and severe trophic skin disturbances of the lower legs and hands with oedema and ulcers, suggestive of autonomic neuropathy. No orthostatic hypotension was observed.

Quantitative sensory testing showed markedly increased or undetectable thermal thresholds for heat and cold sensation in both hands and feet. Dynamic (brush) and static (von Frey hair) mechanical stimuli were not detected. Electrodiagnostic studies revealed absent sensory nerve potentials in the right sural nerve, absent compound muscle action potentials (CMAP) of the right tibial nerve, and markedly reduced CMAP, moderately slowed conduction velocity, and no F waves in the right median nerve. An ECG showed atrial fibrillation. On transthoracic echocardiography there was concentric hypertrophy of the left ventricle, dilatation (51 mm) of the left atrium, no stenoses of the cardiac valves, and normal left ventricular function. The patient had no history of hypertension. Abdominal and thoracic computed tomography detected no tumour mass or lymph node enlargement.

Isoelectric focusing of the serum showed oligoclonal bands identified as IgG λ and κ on immunofixation. In the urine, no Bence-Jones proteinuria was detected, and creatine clearance was within normal limits. The blood leucocyte count was 7.1 × 10^9/μl, 40% of which were lymphocytes. Flow cytometric analysis of the peripheral blood showed that 38% of the lymphocytes were positive for CD19, CD5, CD23, and CD27. These cells showed normal CD20 expression and slight surface expression of λ light chains. A bone marrow biopsy showed multifocal 40% infiltration with

Figure 1  (A) Semithin section of sural nerve stained with methylene blue azure-two, showing massive loss of myelinated fibres and an amyloid plaque (arrow).  (B) Paraffin section with Congo red staining showing birefringence in polarised light.  (C) Cryosection reacted with antibodies to transthyretin (prealbumin Dako, 1:20 000), showing dense immunoreactivity of the plaque for transthyretin.  (D) Cryosection reacted with polyclonal antibodies to human IgG (Dako, 1:1000) showing no immunoreaction on the amyloid plaque (arrows).  (E) Protein analysis by hybrid isoelectric focusing under half denaturing conditions (left), and restriction fragment analysis after amplification of exon 2 of the transthyretin (TTR) gene and digestion with Nsi I for the identification of the ATTR(Val30Met) mutation (right).  Lane 1 represents the patterns of the patient, lane 2 the patterns of a normal control individual, and lane 3 the patterns of a known FAP patient with normal TTR and the ATTR(Val30Met) mutation.
lymphoid B cells of low proliferative activity and no clear birefringence in Congo red staining. A diagnosis of smouldering B cell lymphocytic leukaemia (B-CLL) was made.

Sural nerve biopsy disclosed a dramatic loss of both myelinated and unmyelinated fibres without signs of regeneration (fig 1A). Surprisingly, several amyloid plaques were seen on Congo red staining (fig 1B). Immunohistochemistry showed unequivocal immunoreactivity for transthyretin (TTR) (fig 1C), while polyclonal antibodies to human IgG (fig 1D) stained negative. On skin biopsy of the left lower leg, there was total loss of epidermal nerve fibres. Protein and DNA analysis showed heterozygosity for normal TTR and the amyloidogenic mutation ATTR(Val30Met) (fig 1E) leading to the diagnosis of familial amyloidotic polyneuropathy (FAP).

This patient presents a chance association of B-CLL with “sporadic” FAP. Several cases of a B-CLL associated chronic sensorimotor neuropathy, either caused by neoplastic nerve infiltration or as a paraneoplastic condition, have been described. The initial tentative diagnosis of CLL associated neuropathy was revised when amyloid plaques immunoreactive for TTR were found in the sural nerve biopsy and the ATTR(Val30Met) mutation was demonstrated.

In most cases FAP is caused by a point mutation in the TTR gene. About 80 different mutations of the TTR gene have been identified, the Val30Met mutation being by far the most common. In Europe, this mutation clusters in distinct areas of Portugal and Sweden. Smaller or single families/cases have been described in most other European countries. In Germany about half the known FAP patients are carriers of the ATTR(Val30Met) mutation. Age of onset and penetrance of ATTR(Val30Met) amyloidosis vary considerably. While Portuguese patients from the focus Povoa do Varzim/Vila do Conde develop the disease at a mean age of 31 years, the age of onset among Swedish patients is approximately 57 years. Penetrance is high and progression is rapid in Portugal, but penetrance is low and progression slow in Sweden. Additional genetic and environmental factors probably influence the wide range of both age at onset and severity of FAP. Most FAP patients present with fibre length dependent sensorimotor and autonomic neuropathy. Cardiac involvement, as observed in our patient, is less common and seen in cases of severe polyneuropathy only. Renal involvement is much less prevalent in FAP than in AL amyloidosis, and macroglossia does not occur in FAP. Differentiation of amyloid in tissues by immunohistochemistry is essential for identifying the major amyloidogenic protein. Finally, the diagnosis of FAP must be based on molecular protein/DNA analysis. Although FAP is a disease of autosomal dominant inheritance, a negative family history of polyneuropathy or amyloidosis does not rule out the disease, owing to incomplete penetration or a new mutation. FAP should be considered in all cases of sporadic neuropathy with prominent autonomic symptoms, trophic ulcers, or weight loss, even in countries with a low incidence like Germany. Possibly the prevalence of FAP is underestimated in such countries because of incomplete diagnostic workup.

The recognition of FAP is important for two main reasons. First, treatment is possible by liver transplantation when performed early in the course of disease. Second, diagnosis of FAP in the propositus is essential for identifying relatives at risk for the disease and for providing adequate genetic counselling.

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Aneurysm of the extracranial internal carotid artery presenting as the syndrome of glossopharyngeal pain and syncope

The syndrome of glossopharyngeal pain and/or syncope mimicking idiopathic glossopharyngeal neuralgia has been reported to be associated with a variety of intracranial or extracranial conditions including mass lesions in the parapharyngeal space, the elongated styloid process, and multiple sclerosis. However, aneurysm of the cervical portion of the internal carotid artery (ICA) presenting as episodic glossopharyngeal pain and syncope has not been reported previously to the best of our knowledge. We report here the first such case that was successfully treated by surgical resection of the aneurysm.

A 66 year old woman with a two year history of paroxysmal attacks of pharyngeal pain with occasional syncopal episodes was admitted to our hospital. She had been in good health until two years previously, when she first noticed pain in the region of the left
rhythm, sometimes with radiation to the ipsilateral ear and submandibular area. The pain was neither stabbing nor triggered by swallowing and eating. The attacks recurred about 10 times a day and lasted from several seconds to five minutes. During the year before admission, the attacks had become more frequent, and the duration of pain increased to 30 minutes. She had also felt foreign body sensations in her throat regardless of painful episodes. During the four months before admission, the painful episodes were occasionally accompanied by abdominal pain, diarrhea, and syncope. The episodes of syncope recurred about twice a month. She had been cooled to 34°C and treated with ticlid, a platelet aggregation inhibitor, and at the time of admission to hospital she was having several episodes daily, each lasting up to 15 minutes. She was therapeutically heparinised, her blood pressure was augmented to 150–180 mm Hg with intra venous vasopressors, and florinef treatment was begun to expand her vascular volume. The transient ischaemic attacks persisted unabated despite three weeks of this regimen in the intensive care unit.

Preoperative T2 weighted and magnetic resonance images revealed a few small scattered subcortical signal changes, but no evidence of a previous cerebrovascular event. Cerebral angiography showed severe right supraciloid internal carotid artery and proximal right M1 stenoses with contributions to filling of the middle cerebral tree from pial collaterals. The internal carotid arteries were normal. Transcranial Doppler studies with carbon dioxide vasodilatation challenge showed flow limiting stenosis with compromised cerebrovascular reserve.

On the basis of these data and the refractory nature of the symptoms, we decided to proceed with a high flow revascularisation procedure. An autologous saphenous vein graft was inserted into a proximal M2 branch and the external cerebral carotid artery in end to side fashion. During this time the patient’s systolic blood pressure was maintained between 160 and 180 mm Hg and she had been cooled to 34°C and treated with 20% mannitol (100 g). An intraoperative angiogram, performed immediately after completion of the anastomosis, confirmed patency of the bypass graft and normal filling of the middle cerebral tree. As we were preparing to close the craniotomy, approximately 45 minutes after flow was established through the bypass graft, the brain became massively swollen. A temporary clip was placed across the graft, and the brain relaxed, becoming soft with return of pulsatility. A second intraoperative angiogram showed evidence of hyperperfusion syndrome with the presence of a dense vascular blush throughout the middle cerebral tree (fig 1). Attempts to limit flow through the graft failed and a permanent clip was placed across the graft. Postoperative computed tomography showed intracranial haemorrhage and cerebral oedema with progressive mass effect, commensurate with a worsening neurological state. After discussions with the family, aggressive supportive measures were withdrawn and the patient died.

Symptoms of intracranial stenotic lesions are usually haemodynamic iniology and develop after the collateral blood supply fails
to support metabolic demands despite maxi-
mum oxygen extraction. Management of pa-
tients refractory to treatment with an-
tipilate or anticoagulant agents, blood pres-
sure augmentation, and blood volume expan-
dition is difficult. There is a high risk of stroke, and several studies have argued strongly in
favour of EC-IC bypass in these patients. Con-
sequently, a decision was made in this case to
carry out a high flow bypass from the external
carotid artery to an M2 branch of the middle
cerebral artery. The long nature of the lesion
in the supracallosal carotid, extending into the
proximal M1 segment, was judged not ame-
able to angioplasty. We failed to consider seri-
ously the use of a low flow bypass graft.
Hyperperfusion following cerebral recircu-
larisation is well recognised, particularly in
the context of carotid endarterectomy. Its
occurrence secondary to EC-IC bypass is not
surprising. Intraoperative haemorrhages may
complicate both superficial temporal artery
to middle cerebral artery (STA-MCA) and saphe-
nous vein EC-IC bypasses for carotid occlusive
disease, but evidence for hyperperfusion as
the underlying cause of these haemorrhages
is not established. A dense perfusion blur on
the intraoperative angiogram documented
hyperperfusion following connection of the
EC-IC bypass in this case (fig 1). In chroni-
cally ischaemic brain, hyperperfusion syn-
drome is thought to arise as a result of
disturbed cerebral autoregulation.4 The vascu-
larly is maximally dilated and following re-
toration of flow vascular reactivity is impaired
and unable to vasoconstrict appropriately, to
protect the capillary bed against the increased
perfusion pressure. Low signal oedematous
changes and cerebral haemorrhage result, as
was evidenced on postoperative computed
tomography in this case.

Several factors led to the development of
hyperperfusion syndrome in our case. Preop-
eratively we failed to recognise that altered
microvascular permeability of the chronically
hyperperfused brain may lower the safe
threshold for ischaemia and hypertension.5
Despite neuroprotective measures, temporary
occlusion of the M2 branch for the bypass
may have resulted in a serious degree of
ischaemia, and our routine practice of mod-
estly raising the patient’s blood pressure
during temporary clamping may have been detri-
mental in this setting. Given the increased
susceptibility of the chronically ischaemic
blood brain–barrier to injury, the choice of
a high flow bypass was in retrospect an error.
Depending on the diameter of the graft, saphenous vein bypass grafts provide flows to
110 ml/min.6 By comparison, STA-MCA by-
passes generate flows through the graft in the
range of 10–25 ml/100 g/min.7 Revascularisation is instituted to rescue tissue with blood
flows of 20–25 ml/100 g/min from dropping to
10–15 ml/100 g/min leading to cell death.8 This incremental augmentation of blood flow can be achieved by a low flow STA-
MCA bypass with a lower risk of complica-
tions. Over time, adaptation will occur, allow-
ing increased flow through the bypass graft
to be tolerated and assume a new normal.9 Than if an ischaemic brain is subjected acutely to these high flows.

Patients with intracranial stenosis and
compromised cerebrovascular reserve—high-
lighted as a group most likely to benefit from
EC-IC bypass—are at high risk for hyperper-
fusion syndrome. Strict attention to judicious
choice of bypass conduit, minimising ischa-
emic time for the anastomosis, optimal neuro-
protection, and meticulous control of blood
pressure and intravenous fluids is mandatory
to minimise complications of hyperperfusion
syndrome and optimise the potential benefit
of EC-IC bypass in these patients.

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A Japanese case of steroid responsive myopathy with deficient chondroitin sulphate

In 1998, Al-Lozi et al described a case of steroid responsive myopathy with deficient chondroi-
tin sulphate C that had not been reported before.1 The patient developed diffuse bulbar and systemic weakness with respiratory fail-
ure. While muscle biopsy showed only a mod-
erate degree of type 2 atrophy with one small
perimysial, perivascular mononuclear cell in-
filtration, immunocytochemistry showed an
absence of chondroitin sulphate C in the
endomysium. Prednisone treatment resulted in
a marked increase in muscle strength. Here we
report another case of this interesting and
treatable muscle disorder.

Case history
A Japanese woman first developed neck mus-
cle weakness, dysphagia, and weight loss at
the age of 42 years. She was diagnosed as
having anorexia nervosa at the age of 19 and
received antidepressive drug treatment. Her
symptoms worsened and after a few months she
developed weakness in all of her limbs, sluggish speech, and diplopia, and began hav-
ing difficulty with breathing. She presented in
our outpatient clinic in 1993 at the age of 43.

Her personal and family histories were
negative for neuromuscular disorders. On
physical examination, she was very thin
(162 cm, weight 32 kg), and her temperature was 38.1°C. Coarse crackles were
heard in the right lower lung. Neuromuscular
examination revealed dysarthria, dysphagia,
ataxia, ptosis, limb dysmetria, and weakness
in all directions. Motor and sensory nerve
examinations, and electromyographic studies
failed to show any abnormalities. The tension
was not increased.

Because she was clinically diagnosed as
having atypical occlusory myopathy with respira-
tory failure, a biopsy was performed on the
right rectus femoris muscle. On histochemical
examination, a moderate degree of type 2 atrophy was identified (fig 1A). Immunohistochemical
investigation using antibodies to CD3, CD4, CD8, CD68, and CD22 (Dako Denmark; catalogue Nos 054, 105, 036,
and 093) failed to show the presence of inflam-
matory cells (data not shown).2 At this time, the origin of the patient’s disorder remained
determined.

The patient was given prednisone orally in a
dose of 60 mg/day and her symptoms mark-
edly improved. By day 7, the diplopia, dys-
phagia, and dysphonia had resolved. After four
months, she was discharged with no remain-
ing neurological deficit. Her prednisone dos-
age was gradually reduced and eight years
later, at the time of writing, she was taking a
maintenance dose of 20 mg every other day
and showed no neurological deterioration.

Because the clinical features of steroid
responsive myopathy with deficient chon-
droitin sulphate C—first reported in 1998—were similar to those in our case, we carried out an
immunohistochemical study using a mono-
clonal antibody to chondroitin sulphate

Figure 1 Intraoperative right carotid artery
angiogram showing hyperperfusion, as
evidenced by a dense vascular blush in the
middle cerebral artery distribution.

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(Sigma-Aldrich USA; catalogue No C8035) on frozen muscle samples obtained in 1993 before the patient was started on prednisone. The results showed that there was no binding of antibody to the endomysium or small vessels in the patient’s muscle (fig 1B). In diseased control muscles, diagnosed as motor neurone disease in 1993, chondroitin sulphate staining was demonstrable (fig 1C).

**Comment**

Our patient showed the following clinical features: progressive bulbar symptoms, extraocular movement limitation, and proximal muscle weakness with respiratory failure; steroid responsiveness; and type 2 muscle fibre atrophy with no staining of chondroitin sulphate on histochemical examination. Apart from the limitation of her extraocular movements, the EMG results and the absence of an inflammatory infiltrate on histochemical examination, all other clinical features were markedly similar to those seen in the only other reported case.1 As the antibody used in our immunohistochemical analysis could not discriminate between subtypes of chondroitin sulphate, we were unable to conclude that our patient’s deficient subtype was definitely type C, though the clinical similarities with the other reported case suggest that this was the case.

Type 2 muscle fibre atrophy has been reported in the muscles of malnourished patients.1 Our patient was certainly malnourished as a result of severe bulbar involvement.

The other patient with steroid responsive myopathy was similarly reported to have been malnourished, for a period of 18 months. Thus it is likely that the type 2 fibre atrophy shown in both these patients was caused by malnutrition.

Chondroitin sulphates are major constituents of the extracellular matrix of skeletal muscle and play an important role in binding cytokines as well as in cellular adhesion, differentiation, and signal transduction.2 Thus their disruption in muscle is thought to be involved in the pathogenesis of disease. Al-Lozi et al suggested that myopathy might be caused by a deficit in chondroitin sulphate C developing as a result of immune mediated mechanisms.3 In our case, there was dramatic improvement with steroid treatment, further supporting a role of the immune system in the disease pathogenesis.

In conclusion, we presented a case of steroid responsive myopathy with deficient chondroitin sulphate. This condition should be considered in cases of atypical oculopharyngeal myopathy of unknown origin.

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**Figure 1** (A) Muscle biopsy from our patient showing a predominant type 2 fibre atrophy (adenosine triphosphatase 10.4 staining, ×50). (B) Immunohistochemical staining of muscle obtained from our patient and (C) from disease control subjects, using a chondroitin sulphate specific antibody (×60). Chondroitin sulphate staining was absent from the muscle of our patient.

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**Fulminant progression of hyperammonaemic encephalopathy after treatment with valproate in a patient with ureterosigmoidostomy**

In the absence of liver disease, hyperammonaemia is often considered in the differential diagnosis of encephalopathy and, therefore, the diagnosis of hyperammonaemic encephalopathy may be delayed. We report a case of fulminant progression of hyperammonaemic encephalopathy after valproate treatment in a patient with ureterosigmoidostomy.

A 31 year old patient was admitted because of confusion and agitation. Ureterosigmoidostomy for congenital blind loop oesophagitis was performed when he was 9 years old. Over the past 10 years he was repeatedly hospitalised with episodes of abnormal behaviour, which were blamed on his abuse of various illegal drugs. At this admission, he was disorientated, with lapses into somnolence as well as agitation. Apart from that, his general and neurological status was unremarkable. C-reactive protein was 80.5 mg/l, leucocytes 16160/µl, and blood urea 12.6 mmol/l. Venous ammonia concentration was mildly increased (63 µmol/l). Arterial blood gas analysis showed respiratory alkalosis. All other laboratory analyses including toxicological screening, haematological, biochemical, and microbiological analysis, and cranial computed tomography were unremarkable. Plasma amino acids, organic acids, and orotic acid in urine were normal. After treatment with fluids and amoxicillin with clavulanate for urinary infection, C-reactive protein, leucocytes, blood urea, and sodium concentrations normalised. Soon after admission, he developed generalised tonic–clonic seizures. Anticonvulsant therapy was introduced with 900 mg valproate intravenously followed by continuous infusion (2000 mg/day). Because of repeated seizures, phenytoin was added two days later (1000 mg bolus followed by 1000 mg/day). Despite the antiepileptic treatment, the patient developed a status epilepticus and had to be intubated. The seizures finally stopped after barbiturate coma with thiopental (2–5 mg/kg; 60 000 mg/day) was induced. Arterial blood ammonia concentration was now massively increased (2875 µmol/l). Emergency dialysis was started, and paromomycin and lactulose were administered. Under dialysis, the blood ammonia concentration decreased to 812 µmol/l. However, the patient developed dilated, areactive pupils. Computed tomography showed diffuse brain oedema. The patient died a few hours later, five days after admission. Total plasma carnitine (6 µmol/l, normal range 33–77 µmol/l) and free carnitine (3 µmol/l, normal range 25–54 µmol/l) concentrations were decreased. Acylcarnitine profile also showed decreased but no abnormal individual acylcarnitine species. Repeated analyses of amino acids in plasma and urine, urinary organic acids, and orotic acid showed no significant abnormalities. On postmortem examination, the liver was normal. In both kidneys, multiple abscesses were found. There was massive cerebral oedema and cerebellar herniation.

Ureterosigmoidostomy has been repeatedly associated with episodic hyperammonaemic encephalopathy.1 In this condition, the urine is excreted directly into the sigmoid colon and then excreted during defaecation. Frequent complications include recurrent pylonephritis, faecal incontinence, intestinal malabsorption, and hyperchloreaemic acidosis.1 Hyperammonaemia develops as a result of the increased production of ammonia in the colon from bacterial ureolysis and subsequent absorption of ammonia in the colon. The bowel wall is much more permeable to ammonia than the bladder mucosa, allowing rapid absorption of the ammonia produced within the sigmoid colon. This exceeds the liver’s capacity to detoxify ammonia by the urea cycle. In addition, a proportion of the reabsorbed ammonia is lost from the sigmoid colon throughoesophageal and haemorrhoidal veins. Thus, hyperammonaemic encephalopathy can develop even if liver function is normal.

In the present patient, the diagnosis of hyperammonaemic encephalopathy was not established until the ammonia concentration had risen greatly and the patient was already in critical condition. Urea cycle disorders and organic acidurias were excluded. The laboratory analysis and postmortem examination did not find any signs of hepatic dysfunction. There were no signs of inborn systemic carnitine deficiency. The ammonia concentration increased massively after the introduction of valproate. This can hardly be explained as a
consequence of ureterosigmoidostomy: alone ammonia concentrations as high as these have never been observed in patients with hyperammonaemia following ureterosigmoidostomy. The patient had been treated a few weeks before for urosepsis following cystoscopy and urinary diversion. In our patient, pre-existing episodic hyperammonaemia with encephalopathy and seizures was obviously severely aggravated by valproate induced hyperammonaemia and depletion of carnitine. Patients with valproate induced hyperammonaemia have been repeatedly described, the pathophysiology of which seems to be heterogeneous. In some patients, previously subclinical urea cycle defects have become manifest after treatment with valproate. However, these disorders could be ruled out in our patient. Hyperammonaemia is a frequent side effect of valproate treatment and is often asymptomatic. It seems to occur more frequently in children but is also common in adults, particularly in the presence of other antiepileptic drugs, as was the case in our patient. The exact mechanism of valproate induced hyperammonaemia is unknown but it may appear independently of hepatotoxicity. Valproate has repeatedly been shown to reduce serum and liver carnitine concentrations, both with and without being associated with hyperammonaemia. While most of these patients were children, some cases in adults have been described. Valproate may reduce carnitine concentrations by forming an ester with carnitine, which is co- excreted with organic acids into the urine, or by altering renal reabsorption of acylcarnitine and free carnitine.

In conclusion, our case shows that valproate may greatly aggravate pre-existing, mild hyperammonaemia. We suggest that valproate should be avoided in patients with even slight hyperammonaemia and normal liver function. Equally, we advise the close monitoring of ammonia and carnitine concentrations in patients with ureterosigmoidostomy, such as the one described here, if valproate cannot be avoided.

Palatal tremor and cognitive decline in neuroferritinopathy

Neuroferritinopathy is a recently described autosomal dominant neurological degenerative disorder associated with iron accumulation, particularly in the basal ganglia. All patients found to date have a single adenine insertion between nucleotides 460 and 461 of the feritin light chain gene. This results in a frame shift and is predicted to cause structural alteration of the polypeptide carboxy terminus. Magnetic resonance imaging of the brain shows iron accumulation, and this has been confirmed pathologically with the detection of numerous iron positive inclusions particularly in the globus pallidus. In spite of this, serum ferritin levels are found to be abnormally low or at the lower end of the normal range. Patients tend to present in mid-life with a movement disorder, characterised by chorea, dystonia, and rigidity. In contrast with Hallervorden-Spatz syndrome, which is also associated with accumulation of brain iron, visual and cognitive function is preserved.

Here, we report a patient with genetically proven neuroferritinopathy in which the clinical features included cognitive decline, palatal tremor. These features extend the phenotype of this condition from those previously reported.

The patient was a 49 year old man who developed lingual and oral dyskinetic movements and a slurring dysarthria at the age of 37. Initially, the movement disorder was partially controlled with high dose anticholinergics but then progressed to involve his limbs. Over the next 10 years, he developed dysphagia, unsteadiness, and cognitive decline, particularly of frontal lobe function. His father, paternal uncle, and paternal grandmother had all developed a movement disorder in middle age. A diagnosis of Huntington’s disease was made at that time and was assumed in the patient until disproved by a negative genetic test result. The family continued to seek diagnostic clarification to enable life planning for the patient’s children.

On examination, he was alert and orientated. He scored 7/10 on a mini mental state examination. Detailed cognitive testing showed particular impairment of non-verbal abstract reasoning, with some word retrieval difficulties. He tended to perseverate, and his cognitive estimates were poor. He exhibited pout, palmonmental, and grasp reflexes. He manifested appreciable oral, lingual, and facial dyskinesias. Eye movements were abnormal, with saccadic intrusion into pursuit and use of head thrust to initiate saccades. He had apraxia of eyelid opening. There was no evidence of a pigmentary retinopathy or Kayser-Fleischer rings, and the optic discs were normal. There was a palatal tremor at a frequency of 1 Hz. Examination of the limbs showed pronounced choreiform movements and dystonic posturing. There was no evidence of a peripheral neuropathy or myoclonus.

The following blood tests were normal or negative: full blood count and film, copper studies, creatine kinase levels, ferritin levels (60 µg/l, normal range 25–350 µg/l), liver function tests, and genetic tests for Huntington’s disease (HD), dentatorubral pallidolysian atrophy, and spinocerebellar ataxia 1–3, 6, and 7. Nerve conduction studies were normal. A muscle biopsy was normal. Cerebrospinal fluid analysis was normal, with no evidence of xanthochromia. Magnetic resonance imaging of the brain (fig 1) showed considerable hyperintensity, with a band of surrounding hypointensity on T2 weighting involving the putamen, pallidum, thalamus, substantia nigra, and dentate nucleus. The cerebral cortex was atrophic. A computed tomography scan showed no evidence of cerebral calcification.

Mindful of the clinical features, ferritin deposits, dominant inheritance, and family origin reported in neuroferritinopathy, further genetic testing confirmed the presence of the same A insertion at position 460–461 of the ferritin light chain gene. Investigation of genotypes at six polymorphic microsatellite markers close to the gene on chromosome 19 showed sharing of one allele at each marker. Thus, it is likely that the patient potentially shares the disease associated haplotype with all affected individuals of the originally described families and is another example of inheritance of an initial founder event. Additional evidence comes from the north-west geographical origin of his family.

The differential diagnosis in this case initially included HD, neuroacanthocytosis, and Wilson’s disease, but these conditions were excluded by appropriate investigations. Atypical Hallervorden-Spatz syndrome was also considered, particularly in the light of the appearances on brain imaging, but this condition is recessively inherited and associated with a defective pantothenate kinase gene.
(PANK2). Interestingly, our patient had significant cognitive impairment and palatal tremor in addition to the movement disorders so far described in patients with neuroferritinopathy. In other neurodegenerative disorders, particularly HD, the causative proteins may be involved in iron metabolism. Thus, cognitive impairment may be predicted to occur in neuroferritinopathy, especially in the presence of a pre-existing hyperkinetic movement disorder.

The development of palatal tremor in our patient deserves further explanation. Palatal tremor (previously known as palatal myoclonus) may be classified as essential or symptomatic. It is thought that palatal tremor arises because of functional disruption in “Mollaret’s triangle”, which consists of the inferior olivary nucleus, red and dentate nuclei. The symptomatic form is usually associated with hypertrophy of the inferior olivary nucleus and may arise from vascular lesions, particularly in the cerebellum. Further evidence for this hypothesis comes from a positron emission tomography study, which showed hypermetabolism in the inferior olivary nucleus. Most patients also have cerebellar ataxia. However, palatal tremor may also occur in other conditions including multiple system atrophy, progressive supranuclear palsy, and Alexander’s disease. As in our case, symptomatic palatal tremor is not usually associated with ear clicking. Presumably, in our patient, iron deposition in the dentate nuclei was responsible for disruption of rubral and olivary pathways.

Ferritin is an iron storage protein and alteration in structure of the carboxy terminus could lead to the release of free iron and excessive oxidative stress. In other conditions, such as haemochromatosis, the use of iron chelators has been advocated as a potentially useful treatment. Results, in the main, have been disappointing. Whether free radical scavengers, such as idebenone, have useful therapeutic value in neuroferritinopathy remains to be seen.

Neuroferritinopathy should be considered in all patients with a hyperkinetic movement disorder, imaging evidence of iron deposition within the brain, and an autosomal dominant family history.

### Cocaine induced hypokalemic periodic paralysis

The use of cocaine has been associated with a number of psychiatric, medical, and neurological complications. This is the second reported case of a patient who suffered three distinct episodes of paralysis after engaging in a cocaine binge.

#### Case report

A 33 year old male horse breeder with no significant medical history was evaluated at the Texas Tech Health Sciences Center after he suffered an abrupt onset of generalized weakness. He reported not being able to walk or lift his arms or legs, much less climb up or get down the stairs of his home. He reported no bowel or bladder dysfunction, loss of sensation, headache, nausea, or vomiting. The patient did report mild chest pain at the time. Ten days before his initial evaluation he had suffered a very similar episode but had not sought medical attention. At the time of his evaluation the patient stated that he would be better in 24–48 hours. A very similar event had occurred five years earlier, for which he was seen in an urgent care facility and discharged home; symptoms resolved after 2–3 days. Records of this first episode were not available, although he reported that potassium supplements were provided at that time. Physical examination found an uncomfortable appearing, slightly dishevelled, unshaven man with no spontaneous motor activity. Vital signs were a pulse of 88 beats/min, respiration 16 breaths/min, and blood pressure 132/94 mm Hg. Neurological evaluation found an awake, alert, and oriented person. Speech and language were normal. Cranial nerves were intact. Motor examination found normal bulk with a reduction in tone. Strength was 2/5 in all major muscle groups with a very mild left upper limb predominance. Neck extensors and flexors were 5/5. Bulbar muscles were spared. No myoedema, myotonia, fasciculations, or other abnormalities were noted. The sensory examination was normal and reflexes were symmetric with no Babinski signs. A complete blood count and comprehensive metabolic panel, including electrolytes, renal function, liver function, blood alcohol concentration, and erythrocytosis were performed. Cardiac enzymes were normal. Neuroimaging of the brain and spinal cord were normal. Forced vital capacity and negative inspiratory fraction were normal. Laboratory investigations showed a blood glucose concentration of 6.6 mmol/l; sodium 141 mmol/l, calcium 2.27 mmol/l, and creatine kinase (CK) 395 IU/l. Acetylhydroline receptor antibodies were drawn at the time of admission and subsequently shown to be in the normal range. Two laboratory investigations were of particular interest. The patient’s potassium concentration was 1.9 mmol/l and urine toxicology screen found the presence of cocaine, cannabinoids, and benzodiazepines.

The patient had initially denied any illicit drug use but later admitted to having engaged in a cocaine binge the previous night and before the previous two episodes of weakness. There was no family history of periodic paralysis or other neuromuscular disorders. Supplemental potassium was provided and the patient’s strength gradually improved with rising concentrations of serum potassium. Nerve conduction studies and electromyography were normal at 48 hours after the onset of symptoms. He was discharged home to an outpatient substance abuse program three days later with almost complete resolution of symptoms. At the time of discharge, the serum potassium concentration was 4.5 mmol/l and the CK concentration declined to 131 IU/l. It is not clear why the use of cocaine led to such severe generalised weakness and hypokalaemia in this patient. Nalluri et al. reported a similar case and suggested that the hypokalaemia was caused by an intracellular shift of potassium secondary to the adrenergic effects of cocaine; a hypoaemic form of periodic paralysis in patients suffering from thyrotoxicosis has also been postulated. In their report, as in this case, the patient responded quickly to potassium supplementation. An alternative mechanism may have been cocaine’s potential effects on potassium channels.1 The increased CK and serum glucose concentrations were felt to be the result of cocaine’s effects.

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### Sulcal abnormalities on brain magnetic resonance imaging in the Guillain–Barré syndrome

The Guillain–Barré syndrome is an immunologically mediated condition affecting the peripheral nervous system. There is evidence that Guillain–Barré syndrome, Miller–Fisher syndrome, and Bickerstaff brain stem encephalitis form a closely related spectrum of disorders.1 Magnetic resonance imaging (MRI) findings can help differentiate these conditions have been well described,2 but intracranial findings are infrequent. We report resolution of sulcal changes on serial MRI of the brain concomitant with clinical recovery in a typical case of Guillain–Barré syndrome.

two months after initial presentation showed administration (fig 1). A repeat MRI one week rated inversion recovery (FLAIR) sequences, in tal sulcal hyperintensities on the fluid attenu- sion, showed left parietal and superior occipi- culture and viral studies were negative. Bacterial concentration, and positive globulin. The patient was initially referred to an orthopaedic surgeon for possible lumbosacral spondylitic disease. Unenhanced MRI scans of the lumbar and thoracic spine showed mild degenerative changes and excluded intrinsic cord abnormalities or external compression. Subsequent enhanced cervical spine MRI scans were also negative. He was then referred for a neurological opinion.

Nerve conduction studies revealed signifi- cantly prolonged distal motor latency (median motor distal latency 6.7 to 9.8 ms; posterior tibial distal latency 7.7 to 9.4 ms) and reduced conduction velocities in the median (40.3 m/s), ulnar (39.6 m/s), and tibial nerves (35.7 m/s). F responses were prolonged (> 37 ms) or absent in all four limbs. Cerebrospinal fluid examination showed absent cells with raised protein of 0.8 g/l, normal glucose concentration, and positive globulin. Bacterial culture and viral studies were negative.

The first MRI of the brain, obtained to taking as demonstrated here.

The pathological features of dermatomyositis are characterised by a decreased number of capillaries per muscle fibre and necrosis of single muscle fibres or clusters of fibres at the periphery of the fasciculi. Muscle fibre regeneration and an increased number of capillaries have been shown in dermatomyositis after intravenous immune globulin treatment, but corticosteroids are still consid- ered to be the first line of therapy. In the clineal setting, the effect of treatment is assessed by muscle strength and creatine kinase (CK) levels. Direct measurement of capillary and muscle fibre status can only be done by repeated muscle biopsies. However, apart from the fact that muscle biopsies are invasive, they are also a static representation of muscle tissue at a fixed time point and at a particular location (selection bias).

This is the first time that NIRS, a non-invasive optical method for the measurement of oxygenation and haemodynamics in muscle tissue, has been used to study the effect of treatment in a patient fulfilling the clinical and histological criteria of definite dermatomyositis. A young woman from Aruba, aged 24 years, presented with subacute erythema of the facial skin and severe proximal muscle weakness (arm muscles: mean Medical Research Council (MRC) grade 3; leg muscles: MRC grade 2). Serum CK lev- els were slightly increased (220 IU/l). Five weeks after the onset of symptoms, treatment with corticosteroids was started in our de- partment at a dose of 60 mg/day (for six weeks), the dose being subsequently tapered. CK levels decreased and muscle strength increased (arm muscles: MRC grade 4; leg muscles: MRC grade 3) in week 12.

Muscle tissue oxygenation as a functional tool in the follow up of dermatomyositis

Near-infrared spectroscopy (NIRS) is a direct, non-invasive optical method for measuring local oxygenation and haemodynamics in muscle tissue. Although measurement of local oxygenation by NIRS has been used for the diagnosis of metabolic myopathies, the tech- nique has not previously been applied to inflammatory myopathies. Dermatomyositis is a muscle disorder characterised by comple- ment mediated capillary necrosis, resulting in ischaemia and hypoperfusion. We have now employed NIRS to study the effect of corticosteroid treatment on haemodynamics in muscle tissue in dermatomyositis.

In the related Miller–Fisher syndrome, MRI changes have been documented in the cranial nerves, spinocerebellar tracts, and pons. We postulate that our MRI findings represent a focal manifestation of a wider immunologi- cally mediated reaction within the subarach- noid cerebrospinal fluid bathed space. This focal sulcal reaction probably represents a local concentration of proteinaceous fluid and correlates well with the CSF findings of high protein levels but an absence of cells. These MRI changes resolved with immunoglobulin treatment and clinical recovery. While MRI changes have been reported in the subcortical regions in demyelinating neuropathies, most probably from focal demyelination, sulcal changes have not been described. Serial MRI studies are a sensitive technique for docu- menting cerebral cortical abnormalities in this condition, even in the asymptomatic set- ting as demonstrated here.

References

of Nijmegen, Netherlands). Using this spectrophotometer, which generates light at 905 nm, it is possible to differentiate between oxyhaemoglobin/myoglobin (O2Hb/O2Mb) and deoxyhaemoglobin/myoglobin (Hb/HMb). The optical fibres were placed on top of the flexor digitorum superficialis muscle in the same location for all the measurements. Data were sampled at 10 Hz.

Quantitative NIRS values for oxygen consumption (mVO2) were calculated by evaluating the rate of decrease in [O2Hb] during arterial occlusion, as previously described.1 Reoxygenation rate (ΔO2Hb) was determined as the rate of initial increase in O2Hb measured over three seconds immediately after cessation of arterial occlusion. Both mVO2 and ΔO2Hb were calculated at rest and following rhythmic isometric handgrip exercise at various work intensities. Each exercise session consisted of one minute of exercise at a contraction rate of 30/min (90% duty cycle), immediately followed by 45 seconds of arterial occlusion for the calculation of mVO2 and ΔO2Hb. Whereas mVO2 is a measure of mitochondrial function at a certain work intensity and is dependent on the vascular capacity of oxygen delivery, ΔO2Hb reflects the initial recovery rate at which deoxygenated haemoglobin/myoglobin are resaturated.1 It is therefore directly related to microvascular function. All measurements were performed at the same absolute work intensities.

Figure 1 shows the effect of corticosteroid treatment measured non-invasively and with relative ease by NIRS, in the patient with severe dermatomyositis. Before treatment was begun, resting mVO2 was slightly higher than in healthy controls (0.19 ± 0.14 ml O2/min/100 g, respectively). However, mVO2 during exercise was about 60% lower than in the controls over the whole range of exercise intensities (fig 1A). After three weeks of treatment, mVO2 had already markedly increased. After seven weeks, mVO2 had increased even further and was now only 25% below that of the controls, and within the normal range at several work intensities. Serum CK levels were normalised, while muscle strength had increased. ΔO2Hb (fig 1B) showed similar results, with slow recovery rates before treatment was begun and an increase over all work intensities at the three week and seven week examinations. ΔO2Hb after seven weeks of treatment exceeded the normal mean value.

As NIRS measures local oxygenation and haemodynamics within the muscle, it can give direct insight into the working microvascular system. ΔO2Hb increased during treatment, indicating an increase in capillary function. As a result of the increased capillary function and a possible regeneration of muscle fibres, muscular oxygen availability increased, enhancing oxidative capacity—as reflected by the increase in local muscle oxygen consumption.

Although a muscle biopsy will remain indispensable for the diagnosis of dermatomyositis, NIRS is an interesting and non-invasive tool for monitoring the effect of treatment non-invasively and with relative ease. While both serum CK levels and muscle strength are indirect measures, and muscle biopsies provide only a static fingerprint of the muscle, NIRS measures local microvascular and mitochondrial function directly in the intact and working physiological setting.

**References**


**Figure 1** Effect of treatment in a patient with severe dermatomyositis. (A) Muscle oxygen consumption (mVO2) and (B) reoxygenation rate (ΔO2Hb) measured non-invasively by near-infrared spectroscopy at rest and after exercise at different levels of maximum voluntary contraction force (MVC). Mean values ± SD are shown for the controls.
100 µg/kg/min. Propofol has been suggested as a possible treatment for refractory status epilepticus, although hard data of its therapeu- tic effect are not yet available.6 Propofol has not been used in this condition before but has been effective in two earlier case reports of severe myoclonus: one patient had chloralose poisoning and one had “encephalopathy.”7 In this condition a catastrophic anoxic-ischaemic injury may have damaged the cerebral cortex, basal ganglia, brain stem, and spinal cord and thus the origin of myoclonus remains undetermined. Propofol may termi- nate myoclonus through enhancement of γ amino butyric acid type A receptor. Further experience is needed, but these case reports indicate that good control can be achieved. Propofol’s additional benefit is that intermit- tent neurological assessment remains reliable after discontinuation of propofol.

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Chronic asymmetrical progressive external ophthalmooplegia with right facial weakness: a unique presentation of mitochondrial myopathy

Because of genetic heteroplasmy, the clinical manifestations of mitochondrialopathies are quite varied. We report an unusual presenta- tion in a patient with asymmetric ophthalmoplegia and unilateral myopathic facial weakness caused by a deletion in mitochondrial DNA.

Case history
The patient, a 42 year old white women, was evaluated in our neurology clinic for double vision and right sided facial weakness in early 2000. She first noticed these symptoms in 2000. She first noticed these symptoms in 1994. At that time, she had isolated weakness of adduction of the right eye and she was felt to have a right internuclear ophthalmoplegia. A demyelinating process was considered initially, but magnetic resonance imaging of the brain on four subsequent occasions (annual scan) did not show any structural lesions to support such a diagnosis. The diplo- dia remained unchanged and an insidious facial weakness developed. During the course of her illness she had extensive investigations, including normal cerebrospinal fluid, anti-nuclear antibodies, erythrocyte sedimentation rate, serum protein electrophoresis, and thyroid profile. Rapid plasma reagent and Lyme’s titre were negative. She received several courses of high dose ster- oids without any improvement.

There had been no substantial change in her general physical condition since 1994. Her past medical history was otherwise unremarkable. She is a computer operator and quite active in sports. She has smoked moder- ately for 30 years. She has no family history of any neurological disorders. There was no diurnal variation of her symptoms and no complaint of dysphonia, dysarthria, or limb weakness.

Neurological examination revealed a pupil- sary sparing ophthalmoplegia without ptosis. Specifically she had bilateral exotropia in primary gaze, with total paralysis of adduction of the right eye and mild weakness of abduction of the left eye. She also had mild paresis of infraction of the right eye. Nystagmus was absent in both horizontal and vertical gaze. She had an infranuclear right facial weakness involving both the orbicularis oculi and the orbicularis oris, without lid synkinesis. The strength in the left facial muscles was normal. Ophthalmoplegic myopathy was considered as a possible diagnostic clue. The remainder of her neurological examina- tion was unremarkable. In summary, this patient had a chronic asymmetrical pupil and lid sparing ophthalmoplegia with binocular diplopia and right facial weakness (fig 1). The patient had several normal MRI scans. Previous attempts at treatment with high dose steroids were unsuccessful and the diag- nosis was uncertain. With no structural lesion to explain her clinical condition, we suspected muscular or neuromuscular pathology. An unusual presentation of oculopharyngeal dys- trophy, ocular myopathies, myasthenia gravis, and mitochondrial myopathies were in the differential diagnoses for this progressive ophthalmoplegia. However, we did not have an explanation for her right facial weakness, which we suspected represented a neu-ropathic process. Further investigations showed normal serum lactate and thyroid profile. The blink reflex was normal, excluding the possibility of a cranial neuropathy. Electromyo- graphy of the right orbicularis oculi and orbicularis oris showed myopathic features; on the left side it was normal. This suggested an asymmetric facial myopathy. There was no decremental response of the compound muscle action potential, excluding a possible neuromuscular transmission defect. Muscle biopsy from the right quadriceps muscle showed ragged red fibres. Cytochrome C oxidase stains revealed an absence of staining in many fibres, which showed hiperreactivity on succinic dehydrogenase staining. Electron- microscopic examination of muscle tissue showed an increase in the number and size of the mitochondria, especially in subsarcolem- nal locations—mainly the abnormal cristae structure and paracrystalline inclu- sions. These findings were diagnostic of mito- chondrial myopathy. Genetic testing of muscle tissue was positive for a deletion in mito- chondrial DNA of about 3.5 kilobases, span- ning the ATPase 6 gene to the ND5 gene.

Discussion
Our patient represents a variant of sporadic progressive external ophthalmoplegia with the mtDNA mutation typical of Kearns–Sayre–Daroff syndrome.2 We were able to find one previous report of this syndrome with a similar clinical presentation.11 It was a unique problem because of its atypical clinical fea- tures, which presented a challenging diagno- sis. This case shows the heterogeneity of the clinical manifestations, course, and tissue involvement in mitochondrial disorders.12 The blink reflex was normal and symmetrical, thus suggesting a non-neuropathic facial weakness, which was confirmed by electro- myography. The presentation of this disorder in our patient shows how supranuclear, brain stem, or neuromuscular, or neuromuscular abnormali- ties may be encountered in patients with mitochondrial disorders, either in isolation or in combination. Mitochondriopathies should be included in the differential diagnosis of progressive asymmetric facial palsies, while asymmetrical myopathic facial weakness should be included in the differential diagnosis of infranuclear facial palsies. Our patient’s genetic study identified a deletion of mito- chondrial DNA in a region that encodes different subunits of the respiratory chain complex. Given the clinical manifestations observed, it is clear that the patient has a sig- nificant degree of tissue heteroplasmy.

We would like to emphasise the diagnostic value of the muscle biopsy in cases with atypical clinical presentation and normal serum lactate levels. Genetic testing in blood may become the initial test of choice, with muscle biopsy as an alternative diagnostic aid.

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Figure 1 Observe the wide palpbral fissure on the right, flattening of the right nasolabial fold, and a right exotropia. There was no ptosis of the upper eyelid.