Nerve involvement by malignant melanoma is uncommon but it is a well described feature of the rare amelanotic spindle cell variant of melanoma, termed desmoplastic malignant melanoma. Nerve involvement may be of three types; local invasion of neural tissue, distant spread along perineurium (as in this case), or actual differentiation into neural tissue. Such tumours are termed neurotropic malignant melanoma but are recognised to be a variant of desmoplastic malignant melanoma. The common embryological origin of melanocytes, fibroblasts, and nerve cells from the neural crest supports speculation that melanocytes have the potential to differentiate into fibrous and neural tissue. Clinically the diagnosis of desmoplastic malignant melanoma is often overlooked because it may develop within a very long standing cutaneous lesion, as in this case, or present as a deep fibrous nodule without prominent overlying brown or black pigmentation. Initial diagnoses in one series included viral wart, basal cell carcinoma, pyogenic granuloma, and sebaceous cyst. Histopathological diagnosis may also be difficult. Absence or subtlety of pigmentation with prominent collagenous stromal tissue in the dermis give the tumours a deceptive benign, non-melanocytic appearance. Overlying epidermal lentiginous hyperplasia or the presence of melanocytic junctional proliferation may provide a valuable clue to the likely diagnosis. These features are sometimes absent and diagnosis often relies on the immunocytochemical demonstration of a melanocytic origin. Metastases may show more obvious features of a classical melanoma or retain the desmoplastic features of the primary lesion.

Since desmoplastic melanoma was first described in 1971 over 200 cases have been reported. In one series of 45 patients cranial nerve involvement occurred in 10. Three patients presented with unilateral facial sensory loss, three presented with altered sensation over the face and a cutaneous lesion, and four developed trigeminal or facial neuropathies with recurrent disease. Centripetal spread of tumour cells along branches of the trigeminal and facial nerves, particularly the maxillary nerve and its branches in the face, orbit, and cavernous sinus, may lead to involvement of other cranial nerves.

Although desmoplastic melanoma accounts for less than 1% of all cases of melanoma, any neural involvement is not always a feature. Cases with early cranial nerve involvement may be referred to neurologists before attention is drawn to a cutaneous lesion or to a history of their removal. The case illustrates the need to establish the nature of cutaneous lesions on the face in patients with cranial neuropathies, particularly where superficial branches of the fifth or seventh cranial nerves are involved at an early stage.

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Propranolol in startle induced epileptic seizures

Startle epilepsy is a rare but severe seizure disorder, most often seen in patients with severe brain damage and neurological or mental handicap, but occasionally in patients without signs of cerebral dysfunction. Epileptic seizures of predominantly tonic, atonic-psychomotor, or tonic-clonic semiology precipitated by sudden unexpected mostly acoustic stimuli are the constituent features of this condition. Frequency of seizures is high, with daily attacks in most patients, and falls are common. Spontaneous seizures also occur.

Treatment is difficult. Carbamazepine and benzodiazepines such as clonazepam have been found to be effective in some patients, whereas valproate seemed to be of less value and phenytoin or phenobarbitone were ineffective.

In the Epilepsy Center, Bethel, 24 adult patients with startle induced seizures have been studied and treated since 1980. Twelve of them received clonazepam and 20 received carbamazepine. As a new approach, propranolol was investigated in 11 patients. This was after experimental studies had shown an accentuation of the startle reflex by adrenergic substances, and some anticonvulsant activity. Kolbinger et al had demonstrated the efficacy of metoprolol in a case of startle epilepsy.

We started treatment with 80 mg propranolol, increasing the dose gradually up to 160 mg per day (in two patients up to 240 mg). With propranolol, one patient had no more startle induced seizures, two patients had a greater than 50% reduction in seizure frequency, and eight showed no improvement. There were no side effects and no development of tolerance. The three successfully treated patients remained on propranolol for 82, 19, and 120 months. In the remaining eight patients the drug was withdrawn after treatment failure was apparent. With clonazepam and carbamazepine, one and three patients remained free from startle induced seizures respectively, and one further patient with each drug showed a reduction of more than 50%. Five of seven patients with an initial response to clonazepam developed complete tolerance within two to 16 weeks.

Mean follow up for all patients who were successfully treated was 55 (range 12-120) months. The overall therapeutic outcome was poor. In only 21% of all patients (five out of 24) was a complete control of startle induced seizures achieved. Clonazepam and carbamazepine did not show better results than propranolol. We conclude that propranolol seems to be an additional and safe and effective drug in the treatment of startle epilepsy with efficacy comparable with the standard regimen, but further experience is required.

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Methyl bromide induced neuropathy: a clinical, neurophysiological, and morphological study

Methyl bromide is a colourless, odourless gas used as a fumigant to disinfect soil, grains, and warehouses. Due to its high volatility, dangerous concentrations can accumulate rapidly in working areas. Inhalation is a main route of exposure. Dependent on dose and duration of exposure, methyl bromide can cause both acute CNS symptoms and chronic peripheral sensorimotor neuropathy. Optic atrophy was also described.

A 23 year old Moroccan man working in Italy as a methyl bromide fumigator in greenhouses was admitted because of painful leg numbness. His personal history was unremarkable and he denied alcohol misuse. He had worked for the previous seven months (eight hours a day), wearing a protection mask irregularly. Neurological examination showed asymmetric lower limb weakness and areflexia, stepping gait, and sensory loss for pin prick, position, and vibration. Upper motor and cranial nerve signs were absent. Mental state was intact. Laboratory tests excluded diabetes, hepatic and collagen diseases, syphilis, tuberculosis, HIV, and Lyme related disease. Vitamin B12 and folic acid concentrations were normal. CSF was normal. Organo phosphate and lead poisoning were excluded. One month after cessation of exposure urinary methyl bromide was 0.01 ppm. A colour vision test (Farnsworth-Munsell 100-hue test) showed defective perception of all colours, especially in the red-green axes. Eye fundi and evoked responses were normal.

Electrophysiology showed a severe motor neuropathy of axonal type confined to the legs. The right peroneal action potential was undetectable from the extensor digitorum brevis muscle. The tibial and left peroneal nerves had reduced motor conduction (2.7 and 3.1 SD from mean control value) and muscle action potential amplitude (0.2 and 1.0 mV). Sensory findings in the sural nerve were normal. An EMG showed denervation and loss of motor units in distal muscles.

Tibialis anterior muscle biopsy showed neurogenic changes. A sural nerve sample was examined. Light microscopy showed that the endoneurial area and total fibre density were within the normal range. Multifocal, scattered loss of myelinated fibres (468 = 5200-9500 in normal subjects) was detected, with a prominent loss of fibres larger than 7 ìm (1321 ì 1650-3300), and relative sparing of the smaller axons. Degenerated fibres were seen. The frequency of clusters fell within the normal range and there was no inflammatory infiltrate. On teasing, 38% of the fibres had signs of segmental remyelination; 15% had shortening of all internodes, suggesting regeneration, whereas the incidence of demyelination multifocally distributed along the fibre length was 8%. Electron microscopy showed ongoing degeneration of both myelinated and unmyelinated fibres. Numerous Schwann cell subunits were devoid of axons. Many nodes of Ranvier showed segregated axonal microtubules (figure).

The patient received B1, B6, and B12 vitamins, folic acid, and amyttriptyline (50 mg/day). When re-examined two months later, muscle strength was normal, whereas limb numbness was still present. Peroneal muscle action potential (0.3 mV) was recordable and there were no signs of denervation. Two coworkers, similarly exposed to methyl bromide, had neither clinical nor electrophysiological signs of neuropathy, despite slightly altered colour vision.

The clinical history and laboratory data of this patient excluded known causes of peripheral neuropathy, suggesting that exposure to methyl bromide was responsible. The neurological time course corresponded with previous descriptions, where symptoms appeared after several months of exposure and the patients mostly recovered six to eight months after cessation of exposure.

It has been proposed that methyl bromide causes a distal axonopathy. We report here morphological evidence for nerve abnormalities after exposure to methyl bromide. Loss of large axons and degeneration of both myelinated and unmyelinated fibres were found in the sural nerve. Spared myelinated and regenerating large fibres might account for the normal amplitude and maximal velocity of the sural action potential.

The presence of segregated microtubules in the nodes of Ranvier suggests that methyl bromide impairs axonal flow. As in most types of axonal degeneration, a methyl bromide may cause degeneration of axons with a dying back-like mechanism, more evident at the distal nodes. This neurotoxic agent may either alter the metabolism of the neuronal perikaryon or interfere with the dynamics of axonal transport. A metabolic mechanism for methyl bromide intoxication has recently been proposed. Based on neuropathological similarities with Wernicke's disease, methyl bromide intoxication has been correlated with an altered glycosylation and pyruvate oxidation.

Our patient and two healthy coworkers showed an acquired type dyschromatopsia. This may be a first sign of optic neuropathy, which precedes electrophysiological abnormalities. We therefore propose to use the Farnsworth-Munsell 100-hue test to screen workers exposed to methyl bromide.

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Anti-D immunoglobulin treatment in chronic inflammatory demyelinating polyneuropathy

Patients with immune thrombocytopenic purpura may have increases in platelet count after treatment. The mechanism of action of intravenous
Propranolol in startle induced epileptic seizures.

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