Bilateral substantia nigra involvement in vaccine-associated poliomyelitis

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Parkinsonism is a complication of some viral encephalitis, and poliovirus infection is a possible risk factor for late development of Parkinson disease (PD). There are several reports of PD arising in adults who had poliomyelitis (PM) as children, and pathologic specimens from fatal PM cases disclosed substantia nigra (SN) damage. In this setting, we report a child with vaccine-associated paralytic poliomyelitis (VAP-PM) presenting bilateral SN involvement on MRI.

Case report. A 3-month-old boy came for evaluation 20 days after his first oral poliovirus vaccine (OPV) because of persistent fever, chills, and irritability. Over 5 days, he developed asymmetric lower limb weakness without discernible sensory loss. In the next week, his arms became weak, and poor suckling was noticed. Soon thereafter, frequent apneic episodes and somnolence appeared, and the child required ventilatory support. He was born healthy from cesarean section, and his mother denied any complication during pregnancy. His familial and past medical histories were otherwise unremarkable. On examination, he had asymmetric flaccid quadriplegia with diffuse areflexia and fasciculations. There was obvious bulbar involvement but no objective sensory dysfunction. Pupillary size and reactivity remained normal throughout. Typical ocular bobbing was identified in the first 2 weeks.

Complete blood counts, creatine kinase, electrolytes, hepatic, renal, and thyroid function tests were normal. Serologies for HIV, viral hepatitis, cytomegalovirus, and toxoplasmosis were negative. CSF disclosed 213 leukocytes/mm³ (90% lymphocytes), 165 mg/dL viral hepatitis, cytomegalovirus, and toxoplasmosis were negative. CSF cultures and immunoassays were all negative, as well as a CSF PCR assay for poliovirus. Subsequently, stool culture grew out vaccine-related poliovirus type III. An extensive investigation ruled out any immune deficiency.

Sural and radial sensory action potential amplitudes and conduction velocities were within normal limits. Motor conduction studies revealed small-amplitude compound muscle action potentials with preserved latencies and velocities for age. Although no voluntary muscle activity could be recorded on needle electromyography, prominent spontaneous activity (fasciculations and positive sharp waves) was found in gastrocnemius, quadriceps, deltoid, and biceps brachii 1 month after the onset of paralysis.

Brain MRI showed bilateral brainstem abnormalities (figure, A through C). MRI of the cervical, thoracic and lumbar spine identified lesions in the region of anterior horn cells (figure, D).

Four weeks later, spontaneous breathing and motor recovery began. He was discharged after 2 months and has been followed for 2 years. His cognitive performance is adequate for age, but motor development has been poor. He is not able to stand unassisted. There is residual hypotonia with clear wasting of left leg and right hand muscles. However, no parkinsonian features are currently identified.

Follow-up MRI was performed after 1 and 2 years. There is progressive spinal cord atrophy. Small T2 hyperintense and T1 hypointense lesions are still identified on SN (figure, E).

Discussion. A febrile illness with meningeal signs, flaccid weakness, and residual paralysis lasting more than 60 days after OPV exposure defines VAP-PM. Besides meeting these criteria, our child disclosed ocular bobbing, somnolence, and central apneic episodes, findings consistent with associated brainstem involvement. As proposed previously, this presentation would indeed have features of encephalitis lethargica, which prompted a more extensive neuroimaging investigation.

SN damage documented on MRI has been rarely reported in viral encephalitis. In a few cases of St. Louis, Japanese B, and West Nile encephalitis, parkinsonism was the presenting feature, and SN was found to be involved, particularly the pars compacta. As VAP-PM is a rare condition, neuroimaging findings are scarcely available, particularly with long-term follow-up and with associated brainstem involvement. Despite extensive SN involvement, it is interesting to note that extrapyramidal manifestations were not identified in this patient. They were possibly overshadowed by peripheral findings, namely, hypotonia and weakness.

In a recent Danish study, history of PM was associated with a twofold increased risk for PD. As SN was shown to be affected in fatal PM cases, direct poliovirus damage would be responsible for this increased risk. In fact, viral-mediated neuronal death would not cause PD per se. It would instead make patients especially prone to the consequences of age-related neuronal loss, in a mechanism resembling the postpolio syndrome. By showing SN involvement in vivo, our report gives further support to this hypothesis. Long-term follow-up of this child may help to clarify this issue.

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Clinical/Scientific Notes

Figure. Brain MRI on admission. (A) Axial T2-weighted image showing bilateral hyperintense lesions in substantia nigra. (B) Axial T1-weighted image showing bilateral hypointense lesions in substantia nigra. (C) Axial T2-weighted image showing small hyperintense foci in dorsal medulla oblongata. Cervical spinal cord MRI on admission. (D) Axial T2-weighted image showing symmetric hyperintensities of ventral horns. Brain MRI 1 year later. (E) Residual hyperintense substantia nigra lesions seen on T2-weighted image.
Ocular neuromyotonia secondary to a cavernous sinus meningioma

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Transient diplopia is a common presenting complaint in ophthalmology and neurology clinics. Ocular neuromyotonia (ONM) is a rare cause of transient diplopia and is diagnosed thorough clinical examination, using specific maneuvers to trigger intermittent spasms. We describe a case of left ONM, secondary to ipsilateral cavernous sinus meningioma.

Case report. A 41-year-old previously well woman had episodes of recurrent binocular diplopia for 1 year. These episodes lasted 10 to 20 seconds, occurred 5 to 10 times a day, and were triggered by specific eye movements. She had paroxysmal electrical discharges in the ophthalmic division of the left trigeminal nerve. Examination showed mild proptosis, slight ptosis, and a mild supraduction deficit of the left eye. Diplopia was elicited only after 15 seconds of sustained extreme gaze (see the video, on the Neurology Web site at www.neurology.org). Sustained up gaze triggered retraction of the left upper eyelid, most marked in down gaze. Following sustained right gaze, on attempted return to the primary position, the left eye stayed in an esotropic position and could not abduct for a few seconds. After sustained down gaze, the left hypotropia increased and supraduction was limited. There was no abnormal eye movement after sustained left gaze. Pupil reactions were normal between and during these episodes. A diagnosis was made of a left oculomotor neuromyotonia with a pupil-sparing partial oculomotor nerve palsy and symptomatic left trigeminal neuralgia. Single-fiber electromyography in the frontalis muscle and distal nerve fibers to the levator palpebrae superioris, the medial rectus, and the inferior rectus muscles. ONM is an intermittent ocular deviation due to spasm of one or more eye muscles innervated by the same nerve. Typically spasm is triggered by sustained extremes of gaze.

ONM has been described in 42 patients and involved the oculomotor nerve in 24 of these. Interestingly, sustained eccentric gaze triggered spasm in only 14 cases. The nerve was paretic in 19 cases. ONM was secondary to radiotherapy in 21 cases. Other rarer reported causes include internal carotid artery aneurysm, basilar artery dolichoectasia, midbrain stroke, Grave disease, Paget disease, alcohol, clivus chordoma operated 3 years earlier, cavernous sinus thrombosis secondary to mucormycosis, and diffuse chronic arachnoiditis secondary to thioridazine toxicity. The etiology was unknown in 10 cases. We report a case of ocular motor nerve neuromyotonia due to meningioma.

Case study. A 41-year-old previously well woman had episodes of recurrent binocular diplopia for 1 year. These episodes lasted 10 to 20 seconds, occurred 5 to 10 times a day, and were triggered by specific eye movements. She had paroxysmal electrical discharges in the ophthalmic division of the left trigeminal nerve. Examination showed mild proptosis, slight ptosis, and a mild supraduction deficit of the left eye. Diplopia was elicited only after 15 seconds of sustained extreme gaze (see the video, on the Neurology Web site at www.neurology.org). Sustained up gaze triggered retraction of the left upper eyelid, most marked in down gaze. Following sustained right gaze, on attempted return to the primary position, the left eye stayed in an esotropic position and could not abduct for a few seconds. After sustained down gaze, the left hypotropia increased and supraduction was limited. There was no abnormal eye movement after sustained left gaze. Pupil reactions were normal between and during these episodes. A diagnosis was made of a left oculomotor neuromyotonia with a pupil-sparing partial oculomotor nerve palsy and symptomatic left trigeminal neuralgia. Single-fiber electromyography in the frontalis muscle and distal nerve fibers to the levator palpebrae superioris, the medial rectus, and the inferior rectus muscles. ONM is an intermittent ocular deviation due to spasm of one or more eye muscles innervated by the same nerve. Typically spasm is triggered by sustained extremes of gaze.

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Current treatment is based on cell membrane–stabilizing drugs. Carbamazepine was reported to be effective in 22 of 25 patients. Phenytoin, gabapentine, and acetazolamide were ineffective. In our case, clonazepam was effective for intermittent symptoms, such as diplopia and pain, but oxcarbazepine and acetazolamide were not. The chronic ocular motor defects also improved after treatment, which has been reported in three previous cases. The mechanism of chronic eye movement defects in ONM is not fully understood. In our case, ptosis and hypotropia could possibly be due to ocular motor nerve compression by the meningioma, but these would be unlikely to improve with drug treatment. Spontaneous improvement of chronic oculomotor defects has been previously reported with benign basal skull or cavernous sinus lesions. We postulate that a functional paresis of the infraductor eye muscles or a chronic spasm of the infraductor eye muscles could give rise to these signs. But only functional paresis could also explain the successful treatment of our patient’s ptosis, which also occurred in another reported patient.

ONM is rare, but probably underdiagnosed. Radiotherapy remains the most frequent cause, but MRI is essential to rule out other pathology, particularly compressive lesions. Medical treatment can be effective and may dramatically improve the symptoms.

Transient downbeat nystagmus from West Nile virus encephalomyelitis

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Most human West Nile virus (WNV) infections are asymptomatic, but a minority of patients develops neuroinvasive disease, including a poliomyelitis-like syndrome. Reported ophthalmologic manifestations include optic neuritis, uveitis, multifocal choroiditis, oculomotor palsies, and opsoclonus. We cared for a patient with WNV who developed downbeat nystagmus and ocular flutter with acute areflexic quadriparesis.

Case report. A 64-year-old previously healthy retired steel mill worker presented after 7 days of leg and arm pain and 4 days of progressive weakness. He reported horizontal binocular diplopia, decreased strength, mild dyspnea, and arm tremor. On admission, temperature was 39.2 °C. He was oriented and followed requests. Visual fields were full. Pupils were equal and reactive. He had a mild esotropia in the primary position, with a partial right abduction deficit with right lateral gaze. Downbeat nystagmus was present in primary gaze, and the amplitude increased in right abduction and left adduction. Pursuit was saccadic in all directions, but particularly in vertical gaze. He had intermittent horizontal ocular flutter (video). The optic discs appeared normal. He had scattered gray choroiretinal lesions consistent with multifocal choroiditis. He had asymmetric bilateral facial weakness. There were prominent tongue fasciculations. He had proximal arm muscle weakness (4–), and mild distal weakness (4+). Fasciculations were evident in chest and upper thigh muscles, and he had a coarse tremor of the arms. Sensation of light touch, vibration, pain, and temperature was normal. Arm reflexes were ++, and knee and ankle jerks were absent. Plantar responses were flexor.

On admission, the 7th day of illness, the serum sodium was 131 milliequivalents/L, the white blood cell count was 9.8×10⁹/L, and the hemoglobin was 11 g/dL. The CSF white blood cell count was 813 cells/μL (85% lymphocytes, 5% neutrophils, 12% monocytes), red blood cell count of 37,200 cells/μL, protein 226 mg/dL, and glucose 53 mg/dL. An MRI of the brain without contrast showed lesions consistent with multifocal cerebral and spinal cord dissemination of WNV. A second lumbar puncture on the 13th day of illness showed a white blood cell count of 38/μL (90% lymphocytes, 5% neutrophils, 5% monocytes), red blood cell count of 60,000/μL, protein 226 mg/dL, and glucose 53 mg/dL. An MRI of the spine demonstrated T2 hyperintensity throughout the cord extending from the cervicomedullary junction to the conus. The patient’s vital capacity declined, and he was electively intubated on the 10th day of illness. He received acyclovir and ceftriaxone. He developed meningismus and complete areflexia 2 weeks later. A lumbar puncture on the 15th day of illness showed a white blood cell count of 38/μL (90% lymphocytes, 5% neutrophils, 5% monocytes), red blood cell count of 60,000/μL, protein 226 mg/dL, and glucose 53 mg/dL. An MRI of the brain showed lesions consistent with multifocal cerebral and spinal cord dissemination of WNV. The patient’s critical care team requested help with his eye movements, and the patient was admitted to the neuro-ophthalmology service.

The patient’s cranial nerves were intact on examination. He had full extraocular movements except for an obvious intermittent left downbeat nystagmus with increased amplitude in left head down and right head up gaze (video). He had intermittent horizontal ocular flutter (video). He had a right upper motor neuron facial nerve palsy. He had mild distal weakness of the lower extremities. Spinal fluid WNV PCR was positive. Serum ELISAs showed an increased IgM titer of 2.28 and IgG titer of 3.0. Antimicrobials were discontinued. The patient was successfully extubated after 3 weeks. The downbeat nystagmus, ocular flutter, and right abduction deficit resolved 3 weeks after the onset of symptoms, and strength had improved to 4 in the upper extremities and 1 in the lower extremities.

Discussion. Our patient with WNV encephalomyelitis had two unusual neuro-ophthalmologic findings: downbeat nystagmus and ocular flutter. Downbeat nystagmus usually results from lesions involving the cervicomedullary region or the cerebellar flocculus. Relative excitation of the anterior semicircular canal pathways is thought to be the primary generator of the nystagmus. In our patient, high...
Orthostatic tremor in monozygotic twins

Maria Fiorello Contarino, MD; Marie Laure Welter, MD; Yves Agid, MD, PhD; and Andreas Hartmann, MD

Orthostatic tremor (OT) is characterized by a feeling of unsteadiness during stance that disappears when sitting or lying, associated with a fine-amplitude leg rippling while standing. Surface electromyography (EMG) recordings show a characteristic 13- to 18-Hz tremor.1 Tremor intensity is slowly progressive, sometimes spreading to other body regions, whereas the interval between standing and tremor triggering decreases over time. OT etiology remains unknown. Although most described cases of OT are sporadic, this disorder could be genetic, as suggested by few familial cases2-6 (see table E-1 on the cases of OT are sporadic, this disorder could be genetic, as suggested by few familial cases2-6 (see table E-1 on the Neurology Web site; go to www.neurology.org). We report two monozygotic twins with OT.

Case report. The patients were two male monozygotic twins. Parents were not consanguineous, came from different villages, and remained healthy until their deaths (ages 76 and 87). Family history was negative for tremor or other neurologic disorders. The patients grew up together, lived in the same village, and worked as farmers until age 55, using agriculture chemical products. At age 68, in our outpatient clinic, they had tremor of both legs, occurring when standing and relieved when sitting or lying. Daily living activities were impaired; queuing, talking, and writing while standing were almost impossible. Clonazepam was soon interrupted by both patients because of fatigue and somnolence.

Disease history and presentation were nearly identical in the two brothers when excluding onset age (age 65 for the first born, age 67 for the second born) and milder tremor intensity in the second born. Since youth, bilateral arm tremor had been noticed by both patients while holding objects, which, however, was not bothersome; thus, onset could not be specified. Bilateral leg tremor was observed when standing and abolished when sitting, lying, or leaning against a wall. Latency from standing to tremor onset was about 30 seconds. Mild bilateral arms tremor was noticed when maintaining posture and holding objects, but not at rest. Neurologic examination was otherwise normal in both subjects, revealing no parkinsonian or cerebellar signs.

Surface EMG recordings of bilateral anterior tibial, soleus, deltoid, extensor, and flexor carpi, masseter, and sternocleidomastoid muscles were performed in both patients while sitting at rest, standing, walking in place, pressing palms against a table with arms outstretched from standing position, and maintaining posture with arms flexed at elbows. An accelerometer was placed on one index. A rhythmic 13-Hz EMG activity, alternating between agonists and antagonists and synchronous between the two sides, was recorded in leg muscles (standing, stance phase of walking) and in arms (pressing against a table) (figure). No tremor was recorded in sternocleidomastoids or masseters during posture or contractions.

Discussion. We describe two monozygotic twins affected by OT in which onset age, clinical characteristics, and neurophysiologic findings were similar and correspond to published criteria for OT diagnosis.1 As both patients presented a mild postural arms tremor, a diagnosis of essential tremor (ET) could be evoked. Indeed, the description of coexistent 6- to 8-Hz postural or action tremor in arms in selected OT patients may suggest that OT is pathogenically related to ET or a task-specific variant of ET.2 However,
the prevalence of family history of essential tremor in OT patients is similar to that of the general population (11.5 to 25%).\textsuperscript{3,5,6} Moreover, in our patients, there was no family history for tremor, nor did the clinical or the neurophysiologic findings support the diagnosis of ET. Finally, when appropriately recorded (while pushing against a table from standing), arm tremor had the 13-Hz frequency characteristic of OT.

As both brothers used potential toxic agents (pesticides), this could suggest an environmental etiology, all the more since most reported OT cases are sporadic.\textsuperscript{3,5,6} Even if no acquired factors have ever been reported in association with OT to date, it is not possible to definitely rule out a role for toxic agents.

A genetic cause might be proposed in both brothers, given that they are monozygotic twins and that this symptom onset was almost simultaneous. Few previously described OT patients reported other OT cases among first-degree relatives\textsuperscript{4} (see table E-1). Neither the family we describe nor other published pedigrees allow definite conclusions about the possible transmission modality. As our patients’ parents were unaffected, a recessive or dominant transmission (with incomplete penetrance) could be hypothesized. However, a “second hit” or multifactorial etiology (gene–gene or gene–environment interaction) can also be envisaged, especially with regard to pesticide exposure.

In conclusion, OT is a potentially debilitating disorder with limited treatment options. Elucidating a potential genetic cause may provide hope for identifying causal therapeutic targets. Therefore, we encourage physicians confronted with OT patients to elicit a careful family history and not dismiss the disease as sporadic.

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Co-occurrence of a cavernous malformation and contralateral moyamoya

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We report a young woman who had an intracerebral hemorrhage in her left caudate head from a ruptured cavernous malformation (CM).\textsuperscript{1} During her workup, we found an asymptomatic, moyamoya-like occlusion of her right middle cerebral artery.\textsuperscript{2}

Case presentation. A 37-year-old Brazilian woman with a history of recurrent frontal headache presented to our emergency room complaining of new, worsening biocipital pain for 1 week with nausea, stiff neck, and intermittent right hand weakness. Detailed history revealed no risk for neurovascular disease. She was normotensive with no neurologic deficits. Non-contrast head CT revealed an area of hyperattenuation at the left caudate head consistent with a 20 mL intracerebral hemorrhage (figure, A, page 1602). CT angiogram of the neck and head revealed neither a vascular lesion nor evidence of active contrast extravasation in the area of the hemorrhage; however, it did reveal, contralateral to the hemorrhage, a complete interruption of the proximal portion of the right middle cerebral artery along with well-developed collateral vessels (figure, B), fulfilling criteria for probable moyamoya disease.\textsuperscript{6}

The patient was admitted to our neurointensive care unit for observation. Digital subtraction angiography (DSA) (figure, D) and MRI (figure, E) failed to reveal the etiology of the patient’s hemorrhage. The presence of heterogeneous signal within the hemorrhage on CT (see the figure, A), along with the fact that the caudate head is an uncommon location of spontaneous intracerebral hemorrhage, led us to suspect an underlying lesion predisposed to bleeding. The absence of an arterial malformation demonstrated on CT angiography or DSA raised our suspicion of a CM. Because the patient remained stable with a non-focal neurologic examination, we discharged her, planning further outpatient workup.

Three days later, she returned to the emergency room complaining of drowsiness and vomiting. CT revealed an enlarged left caudate head hematoma with mass effect, intracerebral hydrocephalus, and subfalcine herniation (figure, C). Two days into her neurointensive care unit stay, she experienced rapid neurologic decompensation, with dilated pupils and extensor posturing. After receiving emergent mannitol therapy and placement of an external ventricular drain, she awoke and followed commands.

The following day, she underwent surgical evacuation of the hematoma. Although we faced an unstable lesion in the patient’s dominant hemisphere, as well as the potential risks posed by the presence of fragile moyamoya vessels contralaterally, the patient’s declining neurologic status compelled us to intervene. We performed a bifrontal craniotomy, using an anterior interhemispheric approach as the most direct path to the lesion. Pathologic examination of the resected specimen confirmed a CM (figure, F). She tolerated extubation the day after surgery, and removal of the external ventricular drain 3 days later. Slowly resolving cognitive impairment complicated her hospital discharge, but after about 1 month at a supervised residential facility, she returned home at her previously normal baseline. Six months after her initial presentation, she returned for a successful superior temporal artery to middle cerebral artery bypass procedure, to reduce the risk of rupture from the moyamoya vessels.

Discussion. We cannot explain whether the CM and contralateral moyamoya disease occurred together in our patient by an associated pathogenic mechanism, or simply by chance. It is not surprising that occlusive vasculopathy in one hemisphere may engender hemodynamic changes in the opposite one; however, CMs do not experience arterial pressures, and a direct physical link is not immediately evident. Perhaps, then, a more complex mechanism is at work. Fibroblast growth factor 2 (FGF2, also termed basic fibroblast growth factor or b-FGF), an autocrine/paracrine chemical that stimulates endothelial cell growth and promotes angiogenesis,\textsuperscript{7} is found in histochemical association with CMs,\textsuperscript{4} with particularly strong expression in the case of a CM that ruptured from the moyamoya vessels.

We hypothesize that FGF2, which is known to promote angiogenesis,\textsuperscript{3} is found in histochemical association with CMs,\textsuperscript{4} with particularly strong expression in the case of a CM that ruptured from the moyamoya vessels.

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Reduced penetrance of intermediate size alleles in spinocerebellar ataxia type 10

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Triplet repeat expansions are the disease-causing mutations in nine dominantly inherited spinocerebellar ataxias (SCAs). In 2000, a new type of dynamic mutation was reported in Mexican patients with SCA and seizures, consisting of a (ATTCT) \_ expansion found in intron 9 of the SCA10 gene; normal alleles have 10 to 29 and pathologic 800 to 4,500 repeats.

Methods. We studied 329 unrelated SCA patients. Ataxia was sometimes associated with other features, such as epilepsy, mental retardation, seizures, paraplegia, or tremor; 290 were Portuguese, 39 were from Brazil. Peripheral blood was collected from patients and their relatives, after written informed consent. The
(ATTCT)$_n$ was amplified by PCR with flanking primers and Southern blot was performed as described elsewhere.$^{2,3}$

Results. The modified PCR analysis for (ATTCT)$_n$ expansion showed that three patients (figure, A and B), from two unrelated Brazilian families, with an admixture of Portuguese and Amerindian ancestry, had a continuous ladder exceeding the product range observed for normal alleles at the SCA10 locus. These patients had first shown a single band after PCR for normal allele sizing (figure, C). Expansion size assessment (figure, D) identified one allele with 400 repeats in Patient II-2 from Family 1, and alleles with 760 and 750 repeat units in Patients I-1 and II-1 from Family 2.

These families, unrelated to Brazilian families previously reported, presented a phenotype of SCA without seizures. In Family 1 (see the figure, A), the proband (II-2) was a 59-year-old woman, who reported gait ataxia since age 50 years. Several cases of alcoholism were reported in her ancestry, though the proband was not alcohol-addicted. On neurologic examination, she had gait ataxia, though she was still able to walk independently, together with mild limb ataxia and dysarthria, Babinski sign, and moderate sensory loss in distal portions of the lower extremities. Nerve conduction studies confirmed the presence of axonal polyneuropathy. Cranial MRI showed cerebellar atrophy. The proband (II-1) of Family 2 (see the figure, A) was a 56-year-old man, with slowly progressive gait ataxia since age 20 years. His mother had symptoms since her second decade of life, though clinical assessment was difficult due to the coexistence of a history of alcoholism, absent in him. On neurologic examination, the proband had gait ataxia, mild dysarthria, limb ataxia and pursuit eye movement impairment, limb fasciculations, and bradykinesia. No sensory deficits were found, but nerve conduction studies detected an axonal polyneuropathy. MRI showed atrophy of the vermis.

The (ATTCT)$_n$ showed instability upon transmission (see the figure). In Family 1, the proband had 400 repeats, whereas her father and two of her sibs, all unaffected on neurologic examination, aged 90, 65, and 56, had alleles of 370 and 360 units; her other asymptomatic sib, aged 49 years, also had an abnormal allele, but its size was not assessed due to lack of DNA. In Family 2, the proband inherited an allele of 750 ATTCTs from his affected mother, who had 760 repeats.

Discussion. We describe two Brazilian families of mixed Portuguese and Amerindian ancestry, with an (ATTCT)$_n$, at the SCA10 locus smaller than 800 units. Patients had repeat sizes of 400 to 760 units, thus lowering the threshold for pathogenesis. Alleles with 360 and 370 repeats showed so far no penetrance in asymptomatic subjects aged of 56, 65, and 90. Together with the finding of a 280 repeats allele in an individual with ataxia whose asymptomatic mother had the same size expansion,$^4$ this suggests a range of reduced penetrance for alleles of 280 to 370 repeats. Information on additional chromosomes with 400 repeats is needed to fully understand its pathogenic role. The risk for the offspring of unaffected parents with intermediate alleles, which show reduced penetrance and meiotic instability, is not negligible and, thus, has to be considered according to their instability rate.
and parental origin. Reduced penetrance alleles have been reported for several repeat disorders, including Huntington’s disease and SCA17.

This expansion was initially described in Mexican families and more recently in Brazilians with admixture Portuguese and Amerindian ancestry. Though we cannot exclude the possibility of a Spanish or Portuguese mutation origin, its complete absence so far in several European populations, including Spanish and Portuguese families, makes it more likely to have arisen in Indian populations from the American continent.

Brazilian SCA10 families are characterized by a cerebellar phenotype without seizures irrespective of the repeat size. Considering the intermediate allele sizes in our families, the clinical phenotype seen in Brazilian patients does not seem to be due solely to repeat length, suggesting the involvement of genetic modifiers.

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References

Correction

Fragile X and company: Finding the right diagnosis

In the Patient Page “Fragile X and company: Finding the right diagnosis” by S.H. Subramony, C.A. Friedrich, and J. Jankowiak (Neurology 2005;65:E3–E4), the second author’s name was misspelled. The author’s name is C.A. Friedrich.

The publisher regrets the error.