Acquired episodic ataxia and dysarthria

Commentary by Joanna C. Jen, MD, PhD

The participation of the cerebellum in various feedback circuits involving sensorimotor pathways ensures smooth and precise voluntary and involuntary movement. The superior cerebellar peduncle carries efferent cerebellar projections to the red nucleus and thalamus, as well as the reticular formation. Ascending fibers project to the cortex, from which corticopontine fibers originate to descend to the pontine nuclei, with postsynaptic pontocerebellar fibers carried by the middle cerebellar peduncle, thus closing an important feedback loop between the cerebellum and the cortex. Another feedback loop consists of projections from the red nucleus to the inferior olive, which projects to the cerebral cortex with the red nucleus, thereby indirectly modulating descending rubrospinal and reticulospinal tracts. It is well known that disruption at different levels of these circuits can lead to ataxia. Demyelinating lesions in multiple sclerosis can cause paroxysmal ataxia (presumably from ephaptic transmission of demyelinated axons), yet reports of ischemic subcortical lesions causing paroxysmal ataxia have been rare.

The report from Matsui et al. describes a patient with a left midbrain infarct who, 6 weeks later, developed paroxysmal ataxia and dysarthria with increasing frequency. The authors observed left parietal hypoperfusion by SPECT, which the authors correlated with frequent spells and that improved (but did not normalize) when the patient became free of symptoms after taking phenytoin. The authors hypothesized that sprouting and ephaptic transmission underlies paroxysmal ataxia in this case.

The case study raises interesting hypotheses of CNS injury and repair, but the exact mechanisms require further elucidation and confirmation. Many questions remain unanswered. Did sprouting recur? If so, where did it occur? What triggers paroxysmal ataxia and why should it respond to phenytoin? Diaschisis in the setting of subcortical infarcts is often asymptomatic, and whether parietal hypometabolism accounts for paroxysmal ataxia is speculative; the technique does not allow for temporal resolution for the brief spells in this patient. Phenytoin, which blocks voltage-gated sodium channels, is known to inhibit sprouting. Whether there may be a reorganization of ion channels zone of ischemic injury to account for altered cell excitability and response to phenytoin brings to mind potentially overlapping mechanisms with congenital episodic ataxia and multiple sclerosis. Two subtypes of episodic ataxia, EA1 and EA2, have been shown to result from defects in ion channels abundantly expressed in the cerebellum. There have been anecdotal reports of responsiveness to anti-epileptics in EA1-associated epilepsy caused by heterozygous mutations in a neuronal voltage-gated potassium channel.

The brief and recurrent nature of the patient’s symptoms in this report is more similar to EA1 than EA2. EA1 symptoms are generally not responsive to medications. EA2 can be dramatically responsive to acetazolamide, and a recent report described three EA2 patients who responded to 4-aminopyridine (4-AP), a blocker of potassium channel. Of note, acetazolamide has been reported to be effective in treating paroxysmal symptoms in multiple sclerosis presumed to arise from ephaptic transmission of partially demyelinated axons. Since demyelination affects the distribution of potassium channels, it may be interesting to look into how MS patients with paroxysmal symptoms may respond to 4-AP.

References