Pearls & Oy-sters: Paroxysmal dysarthria-ataxia syndrome

Acoustic analysis in a case of antiphospholipid syndrome

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Pearls

- Paroxysmal dysarthria-ataxia syndrome (PDA), first described by Parker in 1946, is characterized by paroxysmal and stereotyped repeated daily episodes of sudden ataxic symptoms associated with dysarthric speech lasting from few seconds to minutes.1 During the episodes, patients present with slow speech, irregular articulatory breakdown, dysprosodia, hypernasality, variable pitch and loudness, and prolonged intervals, consistent with perceptual characteristics of ataxic dysarthria.2,3
- PDA is a rare neurologic manifestation of either genetic or acquired conditions.2 The most frequent genetic diseases occurring with PDA are episodic ataxias, a group of dominantly inherited disorders characterized by transient and recurrent episodes of truncal instability and limbs incoordination triggered by exertion or emotional stress.4 Among acquired conditions, PDA has been reported mainly in multiple sclerosis (MS), in other immunomediated diseases, or in ischemic stroke.5–7 The common finding among these diseases is the involvement of cerebellar pathways, specifically the crossed fibers of cerebello-thalamocortical pathway in the lower midbrain. Indeed, most of the reported cases of PDA suggest that the responsible lesion is located in the midbrain, near or in the red nucleus,8 where a lesion frequently reveals with dysarthria.9,10

Oy-sters

- Until now, the pathophysiologic basis of PDA remains unknown, as well as the characterization of dysarthria during PDA.
- We present a case of PDA in a patient with antiphospholipid syndrome (APS) evaluated with an acoustic and perceptual analysis of speech to determine the specific pattern of paroxysmal dysarthria.

A 56-year-old woman was admitted due to onset of psychomotor slowing and writing difficulties described as a loss of writing fluency with irregular sizes and shapes of graphemes. There were no cramps or tremor interfering with the task. Symptoms began after an upper respiratory tract infection. Her medical history was unremarkable except for autoimmune hypothyroidism treated with levothyroxine and one unexplained fetal death at the 12th week of gestation. There was no family history of neurologic disturbances, namely of PDA, stuttering, stammering, or cerebellar ataxia. Neurologic examination at admission was unremarkable. Brain MRI revealed increased signal on T2-weighted and fluid-attenuated inversion recovery (FLAIR) sequences in the left cerebral peduncle and left subthalamic region with peripheral contrast enhancement after gadolinium injection (figure, A). EEG did not show paroxysmal activity.

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Neuropsychological assessment revealed impairment of verbal and spatial memory and of executive functions, with a Mini-Mental State Examination (MMSE) score of 28/30. An extensive laboratory workup was done including hematology, chemistry, renal and liver function, coagulation, neoplastic markers (carcinoembryonic antigen, CA 19.9, CA 15.3, neuron-specific enolase, chromogranin, squamous cell carcinoma antigen), and autoimmunity (antinuclear antibodies, antineutrophil cytoplasmic antibodies, extractable nuclear antigen, rheumatoid factor, anti-β2-glycoprotein and antiphospholipid antibodies, C3/C4/CIC, immunoglobulin G [IgG]–immunoglobulin M–immunoglobulin A). All these tests were within the normal ranges. Serum antibodies against *Borrelia burgdorferi* and *Treponema pallidum*, HIV, hepatitis B and C, angiotensin-converting enzyme, and lymphocyte typing were negative. Serum electrophoresis revealed IgG lambda...
type monoclonal gammopathy. CSF cell count was normal while total CSF proteins were slightly elevated. Isoelectric focusing showed identical oligoclonal bands in CSF and serum (mirror pattern). CSF culture for bacteria and nPCR assays for a wide panel of viruses were all negative. In the suspicion of infective encephalitis, the patient was treated with high-dose IV acyclovir. The patient improved gradually during the following weeks with resolution of the writing difficulties and psychomotor slowing. Brain MRIs performed at 1 and 6 months after the discharge showed a progressive reduction of the signal alterations in the left cerebral peduncle and left subthalamic region.

The patient repeated a neuropsychological assessment that showed a complete recovery of the executive deficits and impairment of verbal and spatial memory with a MMSE of 30/30. After 7 months from the discharge, the patient was readmitted to our department for the reappearance of speech and cognitive difficulties associated with multiple paroxysmal episodes of instability, diffuse muscle stiffness, and impairment of speech articulation, triggered by oropharyngeal effort.

The episodes occurred almost every 10 to 15 minutes with duration of 15–30 seconds. Neurologic examination during the episodes showed oculomotor disturbances characterized by bilateral horizontal nystagmus, slow horizontal and vertical saccades, and saccadic hypermetria. There was dysarthria with facial muscles contraction, bilateral ataxia at nose-index and heel-knee testing, retropulsion without backward falls, and gait impairment with impossibility to perform tandem gait without sensory-motor impairments (video 1). Consciousness was always preserved, and intercritical neurologic examination was normal. A new brain MRI showed on T2-weighted and FLAIR sequences the presence of a paramedian hyperintense lesion of the lower midbrain tegmentum with peripheral contrast enhancement (figure, B). The high frequency of paroxysms allowed their registration and evaluation by 2 speech and language therapists. Evaluations were made at a silent voice conversation intensity (<50 dB of background noise) and recorded using a microphone maintained at 20 cm from the patient’s lips. Speech tasks included reading of a short story and a list of 50 words, counting from 1 to 20, a monologue, sustained phonation, and diadochokinesis with fast syllables repetition (pa-ta-ka).

Patterns and degree of dysarthria were evaluated using perceptual and acoustic analysis. The acoustic analysis was performed using the open source Praat software. We quantitatively assessed speech dimensions using objective acoustic analyses with the following aims: (1) to characterize the type and severity of dysarthria; (2) to determine different patterns comparing speech in normal condition and during episodes of paroxysmal dysarthria; (3) to explore the relationship between speech and clinical manifestations. Evaluation in non-symptomatic condition showed sporadic pneumophonic incoordination and mild interictal irregular diadochokinesis. Dysarthric episodes were characterized by slow speech, strained-strangled voice, probably due to effortful squeezing of voice through glottis, pneumophonic incoordination, phonation on residual air, irregular articulatory breakdown, consonant and vowel distortions, dysprosodia, and variable speech rate. Acoustic analysis showed a significant increase of fundamental frequency (mean pitch) during dysarthria episodes, in particular during spontaneous speech, with a difference of 50 Hz, alteration of diadochokinesis with occasional syllable interruptions, inversion, and prolonged phonemes (figure, D–E), and abnormally slow rate of speech during the spontaneous monologue. EEG recordings between episodes and after sleep deprivation did not show any paroxysmal activity. Standard blood examinations, serum neoplastic and infectious markers, autoimmunity, and erythrocyte sedimentation rate were repeated and were normal except for the detection of serum lupus anticoagulant antibodies. A new lumbar puncture was performed; CSF analysis was unchanged with the persistence of a mild elevation of CSF proteins with a mirror pattern on isoelectric focusing. Serum and CSF onconeural antibodies (anti-Hu, Yo, Ma2, CRMP-5, amphi-physin, and Ri), antibodies directed against cell surface antigens (anti-NMDA, voltage-gated potassium channel), and anti-gangliosides antibodies were negative.

Total body CT imaging revealed mild pericardial effusion. In the hypothesis of an autoimmune encephalitis, the patient was treated with IV immunoglobulin (0.4 mg/kg/d for 5 days) and steroids (methylprednisolone 1 g/d for 5 days). The paroxysmal episodes were considered to be compatible with PDA and treatment with carbamazepine (400 mg/d) was started. In the first month after discharge, an important reduction of PDA attacks was observed. At follow-up examination at 3 months after hospital discharge, the patient did not report paroxysmal episodes and neurologic examination was normal. Speech evaluation showed absence of pneumophonic incoordination with the persistence of mild interictal irregular diadochokinesis. A new brain MRI performed 6 months later showed a reduction of the midbrain lesion (figure, C). Repeated autoimmune laboratory testing revealed high IgG anti-β2-glycoprotein I antibodies in 2 separate samples taken after 3 months’ time span. Considering the history of one unexplained fetal death at the 12th week of gestation, neurologic involvement, and laboratory findings, a diagnosis of APS was made based on the revised Sapporo criteria. Treatment with aspirin was started (100 mg/d) and to date, after 18-month follow-up, the patient has not presented any further episodes.

**Discussion**

PDA is an uncommon and rarely reported manifestation of APS. Cerebral involvement in APS is common and characterized by different clinical manifestations, the most frequent of them being cerebral ischemic events. However, in rare cases, APS could manifest with an MS-like syndrome that may mimic the relapsing-remitting form of MS. Usually PDA is...
Carbamazepine has been previously reported to be effective in MS-related PDA, exerting an action against ephaptic transmission between contiguous fiber tracts. Ephaptic transmission is a consequence of the pathologic transverse spreading of fiber activation within a partially demyelinated lesion. Our case confirms that this effectiveness could be extended also to PDA related to other immune-mediated disease such as APS.

Until now, the characterization of PDA-related dysarthria was just based on perceptual and clinical evaluation; only in one case published in 1976 a quantitative analysis of speech was performed in order to better phenotype the characteristic of paroxysmal dysarthria. Our case offered a rare opportunity to study both normal and dysarthric patterns within the same speaker during a single examination period. We wanted to determine specific patterns comparing speech during normal and abnormal episodes of PAD, through perceptual and acoustic analysis. The finding of increased frequency during the episodes could be consistent with a tensing and lengthening of the vocal folds that might result from laryngeal elevation, and it could not be attributed to prosodic or emotional features. Furthermore, differences in fluency and slower articulation rate between dysarthric and normal speech means that articulatory velocity was reduced in the first one. Our clinic–instrumental results show that paroxysmal dysarthria consists of a combination of spastic and ataxic components; this is consistent with the presence of facial muscles contraction during the episodes, as already reported. We can indeed speculate that the diffuse spreading of fibers activation secondary to the pathologic ephaptic transmission within the demyelinated area is not only confined to the cerebellar pathways in the tegmental area of the midbrain but could also involve the more anterior pyramidial tracts located in the cerebral peduncles. Thus, in some cases the paroxysmal manifestations of this disorder could be a result of an abnormal and transient pathologic dysfunction of both pyramidial and cerebellar pathways. Furthermore, the presence of interictal irregular diadochokineses suggests that the cerebellar component of dysarthria persists between PDA episodes, leading us to assume that cerebellar involvement is not only related to a functional paroxysmal alteration of the red nucleus and the dento-rubro-thalamocortical connections but also with anatomical damage of these structures.

Our clinical and instrumental case study expands the comprehension of this rare neurologic syndrome and supports the hypothesis that PDA pathogenesis should be revisited in favor of a combination of spastic and ataxic components.

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