MRI-negative frontal lobe epilepsy with ipsilateral akinesia and reflex activation

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ABSTRACT – [Case records of Epileptic Disorders. Anatomo-electro-clinical correlations. Case 04-2008] The preoperative assessment and surgical management of children and young adults with MRI-negative, frontal lobe epilepsy is often challenging owing to the semiologic expression being unusual or non-lateralizing. Localization based on functional tests may be non-convergent, further compounding the difficulty of surgical planning. We describe a patient with seizures presenting with early, subjective akinesia of the dominant hand that were at times triggered by repetitive motor tasks. Subdural EEG monitoring revealed seizure-origin in the ipsilateral dorsolateral frontal lobe. After resective surgery the patient has had a seizure free period of 9 months, and is free of deficits. This is the first report of frontal lobe seizure semiology consisting of ipsilateral ictal akinesia. [Published with video sequences]

Key words: childhood epilepsy, frontal lobe epilepsy, epilepsy surgery, presurgical assessment

A 17-year-old, right-handed male presented for management of intractable, partial epilepsy. Seizures had begun at age six years and were characterized by brief periods of unresponsiveness and fumbling hand movements, preceded by an aura of vague cephalic sensations or more rarely, an ill-defined sensation in both hands that began minutes or hours before motor seizure-onset. At onset, he complained of an inability to use his right hand. Awareness was preserved during his seizures, which typically lasted 10-20 seconds, and he was mute throughout. Head turning to the right, with accompanying clonic jerking of the left arm accompanied some seizures. Secondary generalized events occurred infrequently, but ictal falls and injuries were common. Seizure frequency was at least weekly. His longest seizure-free interval was two years. Failed medications included carbamazepine and oxcarbazepine, both maintained at high doses, at the time of surgical intervention. Compliance was demonstrated by therapeutic plasma levels of anticonvulsants. Drug side effects included diplopia and severe fatigue.

The patient had been born at term, by elective C-section, after an unremarkable maternal pregnancy. He had borderline hypothyroidism as a younger child. Developmental milestones were normal and he was an above-average student. There was no family history of seizures. His sister...
has neurofibromatosis type 1. Neurological examination revealed two, small, “café au lait” patches, but no other stigmata of neurofibromatosis. There were no focal neurological abnormalities.

Pre-surgical investigations

Previous EEG studies had revealed independent, left and right central and parietal epileptiform discharges, with nonspecific slowing over a wide bilateral field. However, multiple video-EEG studies between ages 15 and 17 years, localized interictal sharp wave discharges in the right frontal and central regions (F4/Fp2 > C4), with a field incorporating the frontal-central midline (Fz/Cz) (figure 1). There were no independent, left hemispheric discharges. His seizures were induced by alternating rapid movements of the hands on two separate occasions, and he consistently described an inability to initiate movement in his right hand at seizure-onset. The earliest observable clinical change was a sudden head tilt/gaze to the right, at times accompanied by atonia of the upper limbs. Longer events evolved into clonic motor activity of the left arm and versive, left, head turning. Secondary generalized tonic-clonic activity accompanied a minority of seizures after drug withdrawal. Speech arrest occurred at seizure-onset, but fluency returned soon after seizure-offset. He remained fully conscious and remembered words spoken during the seizures. Electrographic ictal onset localized to the right frontal and midline regions (figure 2), with later involvement of the contralateral hemisphere during longer events.

Anatomical and functional neuroimaging

MRI studies at 1.5 and 3 T were negative. Functional MRI revealed left-sided language dominance and normal activation patterns during motor tasks. Ictal SPECT revealed a subtle area of hyperperfusion in the right anterior frontal convexity. Repeat ictal SPECT (figure 3) revealed a more robust area of hyperperfusion in the same location, with subtle, increased perfusion in the surrounding regions. PET imaging (FDG/Illumazanil) revealed focal hypometabolism in the left mesial temporal and inferior frontal regions. A cluster of seizures had occurred immediately prior to scanning (figure 4). A repeat PET in the interictal state revealed bi-parietal hypometabolism.

Neuropsychological assessment

Neuropsychological assessment at age 16 demonstrated normal, full-scale verbal and non-verbal intelligence but a specific disorder in mathematics suggesting dominant parietal lobe dysfunction.

Pre-surgical case conference discussion

The sensation of right arm akinesia was correlated to an ictal EEG onset zone in the ipsilateral right frontocentral region; the right head tilt and left arm clonic activity further suggested right frontal lobe involvement. Seizures induced by alternating movements indicate involvement of anterior frontal motor planning systems.

Figure 1. Interictal sleep EEG recording at age 17 years showing frequent, epileptiform sharp and sharp-slow wave discharges over the right frontocentral region, with wide right hemisphere and midline electrode field.
The presence of bilateral upper limb atonia during some seizures suggested possible bilateral involvement of both motor cortices or unilateral involvement of the supplementary sensorimotor area (SSMA).

The epileptogenic region was hypothesized to involve right premotor cortex and ipsilateral SSMA with secondary involvement of the right motor cortex. Left hemisphere involvement was a concern, given the early, negative, right-hand motor phenomenon, PET findings, and dominant hemisphere dysfunction as suggested by neuropsychological evaluation. However, the hyperperfusion in the right frontal region on ictal SPECT reliably co-localized seizure origin to the region of maximal EEG abnormality.

**Surgical plan and procedure, and outcome**

A 48-contact, subdural grid that included motor cortex was implanted over the right frontal convexity, and an 8-contact strip was placed in the right interhemispheric...
region. A 16-contact grid was located epidurally, over the left frontoparietal convexity (figure 5).

Subdural monitoring revealed continuous epileptiform discharges over the right frontal lobe, with a consistent lead from area 6 on the convexity (figure 6), just anterior to the hand motor cortex identified via electro-cortical stimulation mapping. Epileptiform discharges recorded from the right interhemispheric SSMA region were synchronous,
but usually less prominent than those recorded over the convexity. Infrequent, independent epileptiform activity was noted in the left interhemispheric and perirolandic regions (figure 7).

Clinical seizures were accompanied by ictal onset from area 6 over the right frontal convexity and spread to the interhemispheric region (figure 8). Electrical stimulation during mapping over area 6 on the convexity induced his habitual seizures but none were induced upon stimulation of the SSMA region. Independent ictal sequences were recorded from the left frontal convexity contacts, and correlated with episodes of bilateral, tonic, limb stiffening lasting seconds and occurring in clusters.

A corticectomy was performed involving the right frontal premotor region. The resection plane included the regions of early, subdural, ictal involvement and the SPECT abnormality (figure 9). The interhemispheric region was not resected and motor cortex was spared.

Immediately post-operatively, the patient was emotionally blunted and exhibited a resting tremor of the upper limb that was exacerbated by movement. The emotional blunting disappeared over the subsequent week, although subtle, bilateral, upper extremity tremor has remained. The patient had a cluster of non-typical seizures, following a brief reduction of anticonvulsant medication on the third and fourth post-operative days. He has been seizure-free for nine months following surgery, and is taking fewer anticonvulsant medications. Quality of life has subjectively improved. Neuropathology of the tissue revealed Palmini type 2b cortical dysplasia.

Discussion

Our patient’s ictal ipsilateral hand akinesia is an unusual finding that, to our knowledge, has not been previously reported. Clinically, the patient’s motor paresis closely resembled an ictal limb apraxia in that he had no evidence of neuromuscular weakness during his daily activities, and the immobility occurred only during his seizure while he was awake and aware. The occurrence of inhibitory motor phenomena during seizures and the concept of “negative motor areas” have been previously reviewed (Noachtar and Lüders 2000). Immobility of the upper limb may be produced by focal cortical stimulation of prefrontal cortex anterior to the motor face area, but the response is typically contralateral (Penfield and Jasper 1954, Noachtar and Lüders 2000). Stimulation of the anterior SSMA also generates a negative motor response (Lim et al. 1994). Although purely ipsilateral atonia has not been reported during functional stimulation of SSMA, ipsilateral positive motor responses are well described, reflecting the bilateral output of SSMA to motor pathways (Fried et al. 1991). The speech arrest observed during SSMA seizures is also conceptualized as an akinetic event (Noachtar and Lüders 2000).
Figure 8. Subdural seizure-onset during habitual seizures. Initial fast activity and attenuation over the superior convexity is followed by rapid involvement of interhemispheric region (see video sequence 2).

Figure 9. Overlay of motor cortex as defined by functional mapping (green), ictal onset zone (blue), SPECT hyperperfusion (orange), and final resection (dark-bordered region including pink).
In our patient, the convexity and interhemispheric premotor regions were involved early in the ictal discharge, but electrical stimulation of only the convexity led to habitual seizures leading us to postulate that the lateral premotor region was responsible for the ipsilateral akinesia.

It is of interest that our patient’s seizures were at times provoked by sequenced motor tasks, suggesting an intermittent reflex ictal component. Subdural recording revealed ictal involvement of cortical regions known to subserve the planning and sequencing of ipsilateral and bilateral motor movements (Kraakauer and Ghez 2000). In animal studies, the lateral prefrontal area and SSMA participate in the learning phase of coordinated movements; the lateral prefrontal area being preferentially involved (Kraakauer and Ghez 2000). In our patient, seizures may have been triggered by prefrontal and SSMA activation during the fine motor tasks offered. Lateral frontal and premotor seizures with a reflex component have been reported previously, usually as a response to different somatosensory stimuli (Manford et al. 1996).

Our case also highlights the need for careful clinical interpretation of preoperative investigations and a consideration of the limitations of each test. Regional abnormalities on FDG/flumazenil PET imaging strongly correlate with the epileptogenic region in children with focal seizures (Muzik et al. 2000). Initial PET imaging performed on our patient implicated the left temporal and inferior frontal areas in seizure-origin. However, the study was performed during the immediate post-ictal period when PET findings are often inconsistent (Chugani et al. 1993). Relative hypermetabolism may also be recorded in the hemisphere of ictal onset, falsely suggesting ‘hypometabolism’ in the contralateral hemisphere (Henry and Chugani 2008).

This case demonstrates some of the challenges in the assessment of MRI-negative, frontal lobe epilepsy, particularly in the presence of discordant preoperative investigations. Our patient had an additional level of complexity due to early symptoms ipsilateral to the seizure focus. The resective strategy was guided primarily by ictal onset zone and maximal interictal subdural EEG abnormalities, and included the region of hyperperfusion on ictal SPECT. Although the right SSMA was significantly involved, the independent, potential epileptogenicity over the left dictated a more conservative corticectomy of the convexity alone; the long-term success of this approach in our patient is awaited.

References