Kinesigenic reflex epilepsy associated with a glioma in the lateral peri-rolandic region

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ABSTRACT – A patient with kinesigenic focal motor seizures induced by tongue-jaw movement had a grade III astrocytoma clearly co-localizing with the epileptic network in the appropriate peri-rolandic, motor-sensory, lingual-jaw cortical area. The clinical seizure phenomena were time-locked with the EEG epileptic activity. [Published with video sequences]

Key words: kinesigenic reflex epilepsy, peri-rolandic epileptic network, video/EEG

Reflex epilepsies are characterized by seizures induced by a specific temporally associated sensory trigger. They are classified according to the stimuli that trigger them such as visual patterns, lights, sounds, tastes, odors, and movement associated proprioception (Ferlazzo et al. 2005). Kinesigenic seizures (i.e. seizures induced by movement) produced by chewing/eating were originally described by Ahuja et al. (1988). Rarely have reported cases demonstrated clear EEG-neuroimaging correlation. We report a patient with kinesigenic seizures time-locked to the EEG epileptic activity, having a grade III astrocytoma that co-localized with the EEG epileptic network in the appropriate peri-rolandic, lingual-jaw homuncular cortical area.

Case Study

A 67 year old right handed white male with no significant past medical history presented with a single generalized seizure, and then developed simple partial seizures consisting of tonic and then clonic activity of the tongue and jaw to the right. These partial seizures were occasionally spontaneous but usually elicited by talking, the chewing component of eating, and rapid, side-to-side voluntary tongue movements. Although dysarthria occurred during the partial seizures, aphasia did not. There was no alteration of awareness. The patient had a normal general physical and neurological examination. MRI of the brain revealed a 2 cm diameter mass lesion in the left lateral peri-rolandic cortex (figure 1). Biopsy revealed a grade III astrocytoma that

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was treated with radiation and chemotherapy. Generalized tonic-clonic seizures did not recur after phenytoin initiation, but phenytoin did not effectively control the simple partial seizures. Phenytoin was replaced by levetiracetam to avoid anticonvulsant interaction with the chemotherapeutic agents. The simple partial seizures increased after radiation and chemotherapy were completed. Levetiracetam was replaced with extended release carbamazepine with excellent seizure control.

Video-EEG monitoring revealed rhythmic 10 Hz activity that evolved into rhythmic theta activity with accompanying spikes, correlating with tonic followed by clonic activity of the tongue and jaw (see video sequence). The rhythmic epileptiform discharge localized to the left paracentral head region, maximal at the C3 electrode (figure 1). Occasional theta activity was seen interictally in this area, but no other EEG abnormalities were noted.

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**Figure 1.** MRI T1-weighted SPGR (spoiled gradient echo) sequences co-localizing the mass lesion with the EEG epileptic network. A) Axial. B) Coronal. C) Sagittal. MRI shows a grade III astrocytoma in the left peri-rolandic cortex. EEG is in the Laplacian montage. Sensitivity: 7.0 μV/mm. Paper speed: 30 mm/second. High frequency filter: 70 Hz. Low frequency filter: 1 Hz. 60-Hz filter: off. Epileptiform discharge co-localizes with the mass lesion under the C3 electrode.
Discussion

Kinesigenic seizures are rare and have been reported to be evoked by active or passive movements. In our case, the co-localization of the left lateral peri-rolandic, cortical astrocytoma with the left paracentral epileptiform discharge suggests that local area network disruption by the tumor enabled the cortical hyperexcitability. Further mechanistic evidence is provided by the time-locked nature of the epileptiform discharge to the tonic and then clonic tongue and jaw movements to the right (Lopes da Silva et al. 2003).

With our present data, it is not possible to definitively determine whether the proprioceptive sensory input to the motor area, or the voluntary motor activation via praxis circuitry input induced the cortical motor neuron hyperexcitation resulting in the simple partial seizures. But a monkey model of kinesigenic reflex epilepsy with a passive proprioceptive trigger has been reported (Xue and Ritaccio 2006). In one study of eating epilepsy, 98% appeared triggered only by mastication activity (Koul et al. 1989), and the integrated “automatic” motor activities of talking and chewing, which also induced seizures in our patient, probably bypass the voluntary ideomotor praxis areas. All these factors implicate the proprioceptive circuitry input as being the necessary component of the seizure triggering mechanism in our case.

Eating epilepsies, which involve chewing/swallowing movements and therefore can be categorized as movement induced or kinesigenic seizures, have been infrequently reported, but appear more prevalent in the central Asian population (Ahuja et al. 1988, Remillard et al. 1998, Senevirutane 2005). In the Sri Lanka studies, eating induced epilepsy was the most common type of reflex epilepsy and manifested as both partial and generalized seizures. Although focal and generalized epileptic discharges were recorded interictally, none of these eating epilepsy case series were able to record an actual seizure induced by eating or by voluntary lingual-jaw movement. In contrast to our case, neuroimaging (computerized tomography) did not localize the cortical area primarily involved.

Moreover, previously reported cases not only involved signs of peri-rolandic, suprasylvian dysfunction, but were associated with more diffuse lesions clinically demonstrat-