Psychogenic Myasthenia Gravis

To the Editor: Myasthenia gravis (MG) is an autoimmune neuromuscular disorder mediated by the production of antibodies targeting different epitopes of the postsynaptic part of the neuromuscular junction, leading to use-dependent muscular weakness.1 We report a case of functional neurological symptoms mimicking a relapse of MG in a patient who had achieved clinical remission. It has been long proposed that psychogenic neurologic disorders may be due in part to the habituation of idiosyncratic learned, conditioned past response patterns (e.g., psychogenic seizures in an epileptic patient), but this is, to our knowledge, the first report of such behavior mimicking the active phase of the MG illness.

“Mrs. A,” a 33-year-old, right-handed, single mother of three, presented with a short history of fatigue, ptosis, and mild facial and neck weakness. She also suffered with free-floating anxiety symptoms, had an external locus of control, and documented past history of recurrent depression against the background of childhood sexual abuse. A diagnosis of myasthenia gravis was made on the basis of her clinical picture, positive anti-acetylcholine receptor antibodies, and a thymoma detected on CT of the chest, which was revealed in a subsequent thymectomy to be a microinvasive type B3 thymoma (WHO classification). After the operation, she recovered steadily and was a patient without reemergence of other clinical symptoms.

Although she continued to experience attacks, these considerably decreased in frequency (from several a day to once weekly) and severity, allowing for the gradual discontinuation of steroids without reemergence of MG. Moreover, the attacks worsened with MG, as another disorder on the spectrum of mental syndromes associated with MG. Interestingly, the abnormal regulation of inhibitory interneuronal mechanisms, where cholinergic modulation is thought to play an important part, has been proposed as a potential pathophysiological mechanism behind certain psychogenic disorders.2 Hence, one can speculate that this, along with other overt psychological factors, may have led to increased susceptibility to conversion disorder in our patient.

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References

J Neuropsychiatry Clin Neurosci 23:4, Fall 2011
http://neuro.psychiatryonline.org

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