Case Report

Thyroid-Induced Worsening of Parkinsonian Tremor Resistant to Drugs and Subthalamic Nucleus Deep Brain Stimulation

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1. Introduction

Symptoms of hypothyroidism, for example, locomotor slowness and hypomimia, can be easily overlooked in patients with Parkinson’s disease (PD). On the other hand, another sign of PD tremor is also seen in hyperthyroid state. For these reasons, some authors have looked for a link between PD and abnormalities of the thyroid gland with contradictory results. There is a disagreement, whether hypothyroidism is or is not more prevalent in PD patients [1]. Nevertheless, since parkinsonian tremor is known to be exaggerated by thyrotoxicosis [2, 3], it was assumed that the hyperthyroid state leads to increased catabolism of dopamine in the CNS, resulting in a need for higher doses of antiparkinsonian drugs. But the hyperthyroid state causes enhanced tremor not only in PD, but also in essential tremor [2], bilateral chorea-ballism [4], and tremor in Wilson’s disease (not caused by dopamine depletion).

We report on a patient whose parkinsonian tremor worsened and proved refractory not only to dopaminergic and anticholinergic treatment, but also to bilateral subthalamic deep brain stimulation. Thyrotoxicosis was the hidden cause.

2. Case Report

A 61-year-old woman had been diagnosed 16 years previously to have Parkinson’s disease of equivalent type with asymmetrical rigidity, hypokinesia, resting tremor, and mild postural instability. Because of the progressive worsening of motor and nonmotor fluctuations, resistant to oral medication adjustments, we decided to go for surgical treatment. After successful bilateral subthalamic deep brain stimulation (DBS), she enjoyed an excellent outcome. Twenty-one months after the surgery, however, her asymmetrical resting/postural tremor, particularly affecting the lower extremities, markedly worsened (from 1 to 3; score according to the Unified PD Rating
Tremor consists of involuntary rhythmic movements, oscillations, of one or more parts of the body, depending on the mechanical properties of tissues and central neurogenic oscillations. There are several central generators of oscillation in the nervous system, for example, the cerebellothalamic loop or circuits of the basal ganglia (BG). As all of these nuclei are interconnected, their oscillatory activity may converge at the level of the thalamic ventral nuclei [2]. Tremor is observed in the healthy, but it can also be a symptom of disease.

Both the BG and cerebellothalamicocortical (CTC) circuits are responsible for parkinsonian resting tremor. The depletion of pallidal dopamine (due to a lack of projection from the substantia nigra and retrorubral area) leads to excessive synchronization of neurons and thus to enhanced connectivity between the BG and CTC circuits, which exacerbates the tremor activity (the BG are only a trigger) [5]. The cerebellum moderates this activity that is thought to originate in the ventrolateral thalamus with its connection to the subthalamic nucleus and zona incerta. Since the motor cortex is the only part directly projecting to the spinal motor neurons, it is the site of voluntary movements and probably drives the tremor in PD; however, these two activities do not occur at the same time in PD patients. It has been hypothesized that parkinsonian tremor could be (at least partially) an attempt of the motor cortex to compensate for the hypokinetic state by facilitating movement initiation in hyperexcited cortical circuits. This theory is supported by the following facts: (i) tremor is less responsive to dopaminergic treatment, (ii) it is more obvious in earlier stages of PD and in patients with tremor-dominant PD (here the CTC circuits are not yet damaged and are able to produce tremor), and (iii) tremor occurs a few days after hypokinesia and rigidity in animal models of PD [6].

Thyroid hormones (thyroxine and triiodothyronine) increase the expression of calcium currents as well as beta-adrenergic receptors connected with calcium channels in all excitable cells [7], including the central and peripheral nervous systems. The enhanced neuronal excitability or hypersensitivity in the oscillatory loops possibly exacerbates the tremor. The following facts are well known: first, tremor worsens with stress, emotions, hypoglycaemia, and some stimulating drugs (all hyperadrenergic states); second, symptoms are mitigated with beta-blockers, for example, propranolol. Beta-blockers have an immediate effect since they antagonize the adrenergic receptors, whereas antithyroid drugs act more slowly by reducing beta-receptors after thyroid hormone blood levels fall.

4. Conclusion

A hyperthyroid state can markedly exaggerate all forms of tremor as well as other types of movement disorders. This condition can be masked by other symptoms or simply be overlooked. Therefore, if the tremor in a PD patient gradually worsens and proves to be resistant to the usual anti-parkinsonian (or antitremor) treatment, examine the thyroid gland.

Conflict of Interests

The authors declare that there is no conflict of interests regarding the publication of this paper.

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