New-Onset Tourette Syndrome following Human Growth Hormone Therapy

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Dear Sir,

Tics are described as spontaneous, brief, quick, sudden, recurrent, repetitive, purposeless, and stereotyped movements or noises [1]. Tic disorders are classified as (1) transient tic disorder, (2) chronic motor or chronic vocal tic disorder and (3) Tourette syndrome (TS). Of these, TS is the most severe form presenting with both multiple motor and one or more vocal tics and usually starting in childhood [1].

The use of human growth hormone (hGH) in children and adolescents is generally safe. However, it may cause some side effects in a minority of cases [2, 3]. There is no record of tic disorders being associated with hGH therapy. In this paper, a patient with TS whose motor and vocal tics developed during hGH therapy is presented.

Case Report

A 10-year-old boy was admitted to our hospital with a history of short stature. The family history was unremarkable. His height SDS was –2.5. His bone age was nearly every day but they decreased markedly during sleep. At that time, the intensity of shoulder shrug and facial grimace diminished but did not disappear. Thereupon, risperidone, an antipsychotic drug, was started at a dose of 0.25 mg at bedtime and increased to 0.5 mg the following week. In the 3rd month of this treatment, the tics completely disappeared.

Discussion

Our case was diagnosed as TS based on the criteria defined by American Psychiatric Association’s Diagnostic and Statistical Manual of Mental Disorders [1]. The patient had two motor tics and one vocal tic, which occurred nearly every day for more than 1 uninterrupted year, and which developed before the age of 18 years. Furthermore, tics are not due to the direct effects of a known medication.

The cause of TS is uncertain. However, autoimmunity, genetics, neuroanatomy, and neurochemistry are known to play a role in the pathophysiology of TS [1, 4].
Tics might develop following a group A β-hemolytic streptococcal infection, and the infusion of sera into rats from patients with TS resulted in significant oral stereotypy scores, both of which suggest the presence of autoimmunity mechanism in TS [5]. Because streptococcal autoantibodies, known as an autoimmune cause, are a potential risk factor for TS, they should be investigated in children who suddenly develop tics [6]. Our patient had a negative family history of motor and vocal tics.

Previous MRI studies have demonstrated reduced volumes and abnormal volumetric asymmetries of the caudate, putamen, globus pallidus, and frontal lobe in TS patients [1]. SPECT studies have also shown perfusion differences in the same areas [1]. Photon emission tomography investigations have revealed metabolic rate differences of limbic regions of the cortex and striatum, and cortical regions [1]. Although not all imaging studies have yielded definitive findings, they suggest that affected areas might be implicated in the pathophysiology of TS. In our patient, cranial MRI was normal but SPECT demonstrated hypoperfusion of the left frontotemporal region.

Tics are likely to occur due to dysregulation of neurotransmitters in the above-mentioned brain areas. Some drugs such as dextroamphetamine, pemoline, and methylphenidate, known to increase CNS dopaminergic activity, exacerbate TS symptoms, whereas drugs such as haloperidol that decrease the action of dopamine improve TS symptoms [1]. Selective serotonin reuptake inhibitors successfully treat TS associated with an obsessive-compulsive disorder [1]. Selective α1-adrenergic receptor agonists, such as clonidine and guanfacine, are effective for a number of TS patients [1]. These findings suggest that dopamine, serotonin, and norepinephrine may be implicated in the pathophysiology of TS. Altered transmission of these neurotransmitters has been associated with behavioral disorders including anxiety, depression, suicide, aggression and addiction [7]. These disorders have also been reported in children with GHD. A study comparing 109 GHD children with matched controls reported that disorders such as aggressiveness, social withdrawal, somatic complaints, anxiety-depression, delinquent behavior, attention deficiency and thought difficulty were significantly high in GHD [8]. The same study also demonstrated that 72 patients who completed 3 years of GH therapy had a significant improvement in all symptoms. In addition, GHD in adults was associated with detrimental psychological effects such as somatic pain, irritability, depression and increased tiredness, which improve with hGH therapy [9, 10]. These findings suggest that effects of hGH in CNS are biochemically modulated. Recently, studies performed on rainbow trout showed that peripherally administered hGH increased brain dopaminergic activity [11, 12].

Numerous clinical studies have shown that some drugs can lead to exacerbation of TS symptoms [13–15]. Although the mechanism of action is still being elucidated, it is stated that prolonged exposure to medications may alter receptor functioning as well as synthesis or release of neurotransmitters, which are implicated in the pathophysiology of TS. In our case, tics developed 6 months after hGH therapy was started, and hypoperfusion of the left frontotemporal region was the only pathophysiologic finding suggesting that the patient was susceptible to TS. The development of tics in our case was attributed to the dopaminergic activity of hGH based on TS because when hGH therapy was discontinued, the vocal tic disappeared and the intensity of motor tics was diminished; motor tics completely disappeared after risperidone, an antipsychotic drug, was started. However, the literature does not reveal any tic disorder associated with hGH therapy, which may be due to the fact that other patients receiving hGH therapy have no pathophysiologic findings suggesting TS. In addition, because some tics were slight, they might have been overlooked.

In conclusion, patients receiving hGH therapy and having any pathophysiologic findings suggesting TS should be asked about the presence of tics at each visit.

References