Dystonic Tremor in Adult-onset DYT-KMT2B

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Abstract:
KMT2B-related dystonia (DYT28, DYT-KMT2B) is an inherited dystonia that generally begins in the lower limbs during childhood and evolves into generalized dystonia. We herein report a case of adult-onset DYT28 with dystonic tremor. A 27-year-old woman initially displayed right upper limb and cervical tremors over the course of 1 year. A neurological examination also revealed cervical and lower limb dystonia. Although the disease generally develops during childhood, we diagnosed the woman with DYT28, as genetic testing revealed a mutation in KMT2B. Adult-onset patients with DYT28 might also show uncommon symptoms as well as DYT-TOR1A (DYT1).

Key words: KMT2B-related dystonia, DYT28, dystonic tremor, dystonia

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Introduction

KMT2B-related dystonia (DYT-KMT2B, DYT28) is an inherited dystonia discovered in 2016 (1) that usually develops during childhood. Typically, dystonia symptoms of DYT28 begin in the lower limbs and subsequently progress to generalized dystonia with the inclusion of the upper limbs, pharynx, and trunk (2). Most cases develop during childhood; therefore, the clinical characteristics of adult-onset DYT28 have remained unidentified.

We herein report a Japanese patient with DYT28 whose disease onset began at 26 years old, with upper limb and neck tremors as the initial symptoms.

Case Report

A 27-year-old Japanese woman presented with a 1-year history of tremors in the right upper limb and cervical regions. She had a medical history of anxiety disorder and a mild intellectual disability; in addition, she had a family history of depression (father and sister) and intellectual disability (brother), but no family history of hyperkinetic movement disorders (Fig. 1).

The patient’s height was 140 cm, and she had an elongated face with a bulbous nose. She showed involuntary movement characterized by an irregular tremor in the neck and postural tremor of the right-dominant upper limb and neck. However, resting and task-specific tremors were not observed. The tremor was exacerbated by mental load (e.g., while performing calculations). In addition to tremors, we observed mild generalized dystonia, slight retroflexion, and left rotation in the neck during the appearance of tremors as well as bilateral dorsiﬂexion of the first toe while walking. Furthermore, although we observed right-dominant hypertrophy of the sternocleidomastoid and paraspinal muscles, she had no subjective symptoms of dystonia, except for neck pain.

Her cranial nerve examination, muscle strength, tendon reﬂexes, finger-to-nose test, and diadochokinesia tests reported normal ﬁndings. Furthermore, her laboratory tests revealed no abnormalities, and brain magnetic resonance imaging also revealed no abnormalities, including the basal ganglia. The accumulation on dopamine transporter single-photon emission computed tomography (DAT SPECT) showed a slightly irregular shape, with the right striatum having a specific binding ratio (SBR) of 3.90 and the left striatum a value of 3.83 (Fig. 2). Surface electromyography showed continuous discharges from the bilateral sternocleidomastoid muscles (SCMs) at rest and alternating discharge

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from the right forearm when the patient held her arms in an outstretched posture (Fig. 3). Her intelligence quotient was 70 on the Suzuki-Binet test.

Genetic testing was performed on the patient’s blood-derived DNA, and a new heterozygous nonsynonymous substitution was detected in the \textit{KMT2B} gene c. 5285 G>A (p. Arg1762His) (Fig. 4A). This variant is not present in the GenomAD (https://gnomad.broadinstitute.org/) or human genetic variation databases (https://www.hgvd.genome.med.kyoto-u.ac.jp/) and is not considered to be present in the general population. The affected amino acids are conserved among species (Fig. 4B), and the prediction scores, including Sorting Intolerant From Tolerant, MutationTaster, and Polyphen-2, suggested that this mutation is pathogenic. Based on her characteristics of dystonia, short stature, elongated face with bulbous nose, anxiety, and intellectual impairment, we diagnosed her with DYT28.

She was treated with levodopa/carbidopa (100/10 mg 3 times daily), but her symptoms did not improve. However, on treatment with clonazepam (0.5 mg) and botulinum toxin injection into the bilateral SCMs and trapezius, her tremors and dystonia improved slightly.

**Discussion**

In this case, the onset of DYT28 presented as a dystonic tremor in the upper limb and neck at 26 years old. DYT28 generally shows progressive, generalized dystonia starting in the lower limb region and spreading to the upper limb, cervical, pharynx, and trunk during childhood. DYT28 also exhibits other neurological symptoms, including myoclonus, anxiety, intellectual disability, spasticity, and epilepsy. In recent years, there have been a series of reports on DYT28, which may account for 10-22% of cases of child-onset generalized dystonia, suggesting that it is a more common disease than was previously thought (2, 3). The clinical profile of KMT2B-related disorders is changing, as a psychiatric variant without dystonia symptoms has been reported (4). However, adult-onset cases are rare; adult-onset dystonia is generally less markedly affected by certain genes than childhood-onset dystonia. (5). Two previous cohort studies of DYT28 identified only 1 case each of adult-onset dystonia among 27 and 53 patients, respectively. However, no cases of dystonic tremor were reported in their cohorts (4, 6).

In Table, we summarize all adult-onset cases reported thus far. Interestingly, the onset site was not the lower limbs in any adult-onset cases; in three cases, the onset began in the upper limbs (4, 7, 8); in one case, in the neck (9); in one case, in the larynx (7); and in two cases, the site was unknown (6, 10). Furthermore, only one case other than the present case showed tremor at an onset age of ≥18 years old. These findings suggested that adult-onset DYT28 may present with atypical clinical symptoms. DYT-TORIA (DYT 1) is the most representative inherited dystonia, but there is a difference in symptoms between childhood- and adult-onset cases. Typically, DYT1 has a phenotype that presents in childhood with an onset in the lower limbs and subsequently generalizes. In contrast, patients with adult-onset DYT1 presented with focal dystonia or tremor within the
Figure 3. Surface electromyography. Note the alternating discharges from the right forearm. The patient held her arms in an outstretched posture. SCM: sternocleidomastoid, NE: neck extensor, PSM: paraspinal muscle, BB: biceps brachii, TB: triceps brachii, WF: wrist flexor, WE: wrist extensor.

Figure 4. A: Sanger sequencing results for the KMT2B gene in the patient. An electropherogram of the Sanger sequencing result. C.5285G>A (p. Arg1762His). B: Levels of conservation of the p. Arg1762 domain in KMT2B among different species (references cited in https://genome.ucsc.edu/cgi-bin/hgTracks?db=hg19&hla=KMT2B&domainType=cons&domainTypeDefault&show=0&hide=0&nogenome=0&format=bed&geneline=1%3A36221601%2D36221630&hgsql=1183100549_XpNLOGDzkNOeGN0AIw7srA9kOaX).
family of typical generalized dystonia (11). Similar to DYT1, adult-onset DYT28 cases may show a clinical course different from that of childhood-onset cases; therefore, further analyses are warranted.

The results of DAT SPECT suggested dopaminergic dysfunction in our case. The striatal accumulation was slightly irregular in shape, and the SBR values were low. The lower limit of the SBR in the Japanese database for patients in their 30s is 6.82 (12), suggesting that the SBR in this case was decreased, although the normal value in the 20s is not established. A recent study revealed that patients with generalized dystonia show a mildly reduced DAT SPECT accumulation (13). The results of DAT SPECT in our case were also considered to reflect a disturbance of the dopaminergic system in dystonia.

We obtained written informed consent from the patient. This study was performed in accordance with the Declaration of Helsinki and was approved by the ethics committee of Tokyo Metropolitan Neurological Hospital and Tokyo Women’s Medical University.

The authors state that they have no Conflict of Interest (COI).

References


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