Pediatric-Onset Dystonia Associated with Bilateral Striatal Necrosis and G14459A Mutation in a Korean Family: A Case Report

We describe a Korean family presenting with pediatric-onset, progressive, generalized dystonia with bilateral striatal necrosis and the homoplasmic G14459A mutation in the mitochondrial ND6 gene. The G14459A mutation has been reported in families presenting with Leber hereditary optic neuropathy (LHON) alone, LHON plus dystonia, or pediatric-onset dystonia. The proband had shown dysarthria, progressive generalized dystonia, and spasticity at 5 yr. Brain MRI demonstrated bilateral striatal necrosis. Additional investigation of family members revealed the presence of homoplasmic G14459A mutation in asymptomatic individuals. The clinical manifestation of the homoplasmic G14459A mtDNA mutation within the same family showed asymptomatic or pediatric-onset dystonia, without optic neuropathy. This study reemphasizes that the G14459A mutation is a candidate mutation for maternally inherited dystonia, regardless of optic neuropathy, and supports the hypothesis that nuclear genes may play a role in modifying the clinical expression of mitochondrial disease.

Key Words: Mitochondrial Diseases; Basal Ganglia; Necrosis; Dystonia; Nucleotide Position 14459

INTRODUCTION

Mitochondrial DNA (mtDNA) point mutations have been associated with a wide range of clinical presentations, ranging from pure myopathies to multi-systemic disorders. The G14459A mtDNA mutation changes a moderately conserved alanine residue to a valine within the most evolutionarily conserved region of the ND6 gene, a component of complex I of the mitochondrial electron transport chain (1). The G14459A mutation has been associated with Leber hereditary optic neuropathy (LHON)/pediatric-onset dystonia. The proband had shown dysarthria, progressive generalized dystonia, and spasticity at 5 yr. Brain MRI demonstrated bilateral striatal necrosis. Additional investigation of family members revealed the presence of homoplasmic G14459A mutation in asymptomatic individuals. The clinical manifestation of the homoplasmic G14459A mtDNA mutation within the same family showed asymptomatic or pediatric-onset dystonia, without optic neuropathy. This study reemphasizes that the G14459A mutation is a candidate mutation for maternally inherited dystonia, regardless of optic neuropathy, and supports the hypothesis that nuclear genes may play a role in modifying the clinical expression of mitochondrial disease.

CASE REPORT

A Korean family was enrolled in this study. The proband (individual III-12), a younger sister (III-13), and a younger brother (III-14) were a 34-yr-old man, a 32-yr-old woman, and a 30-yr-old man, respectively, who had non-consanguineous parents (Fig. 1A). Their health was unremarkable until 5 yr of age, when they developed unilateral distal dystonic posture and gait disturbances. The phenotype progressed and they all currently showed severe dysarthria, contractures of both ankle and wrist, both knee contractures, spasticity in both lower extremity, spinal scoliosis, dystonic hands, and generalized hypotonia, retropulsion, with mild mental retar-
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Their clinical features including time of onset were very similar to those of the proband. The proband and younger brother (III-14) was unable to communicate by phone because of severe dysarthria. They could be ambulant by only wheelchair because of multiple joint contractures, spasticity of lower extremity, and dystonia of foot. Vision and hearing were unaffected. Brain MRI showed the presence of bilateral symmetric tissue losses in both the putamen and caudate nucleus (Fig. 2). An increase in lactic acid or amino acids levels in plasma was not observed, and no abnormalities in plasma copper or ceruloplasmin levels were seen. Nerve conduction tests showed no abnormality. The nephew (IV-3), a 6-yr-old boy, exhibited right hand dystonia, right ankle contractures and gait disturbances. However, he was able to walk and swim by himself. He was normally delivered. His onset was at 5 yr old, and it was also similar to that of proband. He was still normal in speech, intelligence, and stature. His MRI revealed bilateral high signal intensity in T2-weighted images, without necrosis (Fig. 2). The maternal uncle (II-3) and maternal cousins (III-1 and III-2) suffered from progressive dystonia. One maternal cousin (III-1) committed suicide in his third decade. The proband’s mother (II-7), sister (III-11), and maternal aunt (II-2) were asymptomatic.

**Genetic analysis**

Six members (II-7, III-11, III-12, III-13, III-14, and IV-3) of the family were investigated by molecular genetic tests (Fig. 1A). Four (III-12, III-13, III-14, and IV-3) had progressive, generalized dystonia. Two (II-7, III-11) were asymptomatic. After obtaining informed consent, genomic DNA was isolated from peripheral blood leukocytes using a Wizard genomic DNA purification kit (Promega, Madison, WI, USA), according to the manufacturer’s protocol. The samples were analyzed for the presence of 15 point mutations associated with dystonia with bilateral striatal necrosis, LHON, or MELAS (A3243G (7), T3271C, T3308C (8), G3460A, A8296G (9), A8344G, T8356C, G8363A, T8851C (10), T8993G (11), T9176C (11), G11778A, G14459A, T14484C, and T14487C (12) by direct sequencing. In addition, the A3203G muta-

![Fig. 1. (A) Pedigree of a Korean family with maternally inherited, pediatric-onset dystonia carrying the G14459A mutation. (B) Direct sequencing analyses of the mitochondrial DNA for the G14459A mutation. Circle, female; square, male; black symbol, affected; dot, asymptomatic carrier; diagonal line, deceased.](image-url)

![Fig. 2. Brain MRI of the proband (A, B) and his nephew (C, D). (A) The T2 weighted image of the proband show bilateral striatal hyperintensities with necrosis. (B) The T1 weighted image of the proband show bilateral hypointensities with necrosis. (C) The T2 weighted image of his nephew shows bilateral putaminal hyperintensities without necrosis. (D) The T1 weighted image of his nephew shows bilateral putaminal hypointensities without necrosis.](image-url)
tion, previously identified in a Japanese family, was analyzed (7). The mitochondrial DNA was amplified by polymerase chain reaction (PCR) using primers designed by the authors (available upon request) and a thermal cycler (Model 9700; Applied Biosystems, Foster City, CA, USA). Direct sequencing was performed using the BigDye Terminator Cycle Sequencing Ready Reaction kit (Applied Biosystems) on an ABI Prism 3100 genetic analyzer (Applied Biosystems). The direct sequencing analysis revealed a homoplasmic G14459A mutation in each of the patients examined (II-7, III-11, III-12, III-13, III-14, and IV-3; Fig. 1B). Thus, two individuals (II-7, III-11) were found to be asymptomatic carriers.

**DISCUSSION**

The clinical features of the patients in this Korean family included progressive, generalized dystonia, multiple joint contracture, and spasticity, which were attributable to striatal degeneration. In previous reports, the G14459A mutation was observed in one Hispanic (2), five Caucasian (3-6), one African-American (3), and one Japanese family (7) (Table 1). The general features of previously reported cases were optic neuropathy, progressive generalized dystonia, mild mental retardation, and spasticity. In our family, pediatric-onset dystonia, spasticity, and dysarthria were the major clinical presentations. Based on the clinical features between Caucasian

**Table 1. Families reported as having maternally inherited dystonia with the G14459A mutation**

| Race of Family | Reported family member | Age at onset | Symptoms | Cognition | Ophthalmopathy | Lesions on neuroimaging | G14459A Mutation type | Phenotype | Reference |
|----------------|------------------------|--------------|----------|-----------|----------------|------------------------|-----------------------|-----------|-----------|
| Hispanic       | Proband (IV-36)        | 2 yr         | Progressive generalized dystonia | Normal | Normal | Bilateral lesions in the putamen and caudate nucleus | Homoplasmic | Pediatric onset dystonia | (2)       |
|                | III-5                  | Not described | Asymptomatic | Not described | Normal | Not described | Homoplasmic | Asymptomatic |           |
|                | III-10                 | 32 yr        | Bilateral optic atrophy and dystonia | Not described | Not described | Present | Not described | Heteroplasmic | LHON      |
|                | IV-25                  | Not described | Optic atrophy | Not described | Present | Not described | Not described | LHON plus dystonia |           |
|                | IV-26                  | Not described | Optic atrophy | Not described | Present | Basal ganglia lesions | Homoplasmic | LHON      |
|                | IV-35                  | 13 yr        | Mild dystonia and intellectual impairment | Delayed | Normal | Not described | Homoplasmic | Pediatric onset dystonia |           |
|                | V-11                   | 5 yr         | Mild generalized dystonia | Not described | Normal | Basal ganglia lesions | Homoplasmic | Pediatric onset dystonia |           |
| African        | Proband                | 42 yr        | Gradual, painless visual loss | Normal | Present | Not described | Heteroplasmic | LHON | (3)       |
|                | Daughter               | 19 yr        | Gradual, painless visual loss | Normal | Present | Unilateral lesion in the right putamen and bilateral lesions in the caudate nucleus | Homoplasmic | LHON |           |
| Caucasian      | Proband                | 34 months    | Generalized dystonia and atherosis, dysarthria | Delayed | Normal | Bilateral extensive lesions in basal ganglia | Heteroplasmic | Pediatric onset dystonia | (3)       |
| Caucasian      | Proband                | 18 yr        | Bilateral subacute visual failure, hearing loss, ataxia | Normal | Present | Lesions in dorsal midbrain and right red nucleus | Heteroplasmic | LHON plus dystonia | (4)       |
| Caucasian      | Proband (IV-10)        | 3 yr         | Stroke, dystonia | Previously delayed | Normal | Bilateral lesions in the putamen | Homoplasmic | Pediatric onset dystonia | (5)       |
|                | III-6                  | 35 yr        | Asymptomatic | Normal | Present | Bilateral and temporal signal abnormalities, possible hematoma | Homoplasmic | Asymptomatic |           |
|                | IV-2                   | 5 yr         | Limp Hemiparesis | Normal | Normal | Bilateral lesions in the putamen | Homoplasmic | Pediatric onset dystonia |           |
|                | IV-8                   | 7 yr         | Cognitive delay, Hemiparesis | Delayed | Normal | Unilateral lesion in right sided putamen | Homoplasmic | Pediatric onset dystonia |           |

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and non-Caucasian in Table 1, racial difference seemed to be not apparent. The most striking difference in our family could be no LHON manifestation such as visual disturbance or optic nerve atrophy. Although delayed manifestation of visual disturbance was reported in Japanese family, which was shown at 38 yr, at least one person of their pedigree had visual disturbance or optic nerve atrophy in previous reports (2-7). Accordingly, we don’t know whether any of our family members will have delayed manifestation of optic neuropathy or not.

Among the described cases (26/27) with the heteroplasmic or homoplasmic G14459A mutation in Table 1, age at onset, dystonia, impaired intelligence, and basal ganglia abnormality in neuroimaging was not statistically significant between heteroplasmic and homoplasmic mutation groups (Table 2). The heteroplasmic G14459A mutation has been associated with heterogeneous clinical phenotypes, varying from asymptomatic to dystonia, LHON, or dystonia plus LHON among family members, while the homoplasmic G14459A mutation has primarily been associated with pediatric-onset dystonia.

Although neither heteroplasmy nor homoplasmy for this mutation is a direct predictor of LHON or a dystonia phenotype, many individuals are essentially homoplasmic for the mutation. Thus, a threshold effect of the mtDNA mutation may contribute to the phenotype of the mitochondrial disease. However, a threshold effect could not explain why some cases with homoplasmic mutation were asymptomatic. Furthermore, the G14459A mtDNA mutation was also

Table 1. (Continued from the previous page) Families reported as having maternally inherited dystonia with the G14459A mutation

| Race of Family | Reported family member | Age at onset | Symptoms | Cognition | Ophthalmopathy | Lesions on neuroimaging | G14459A Mutation type | Phenotype | Reference |
|---------------|------------------------|-------------|----------|-----------|----------------|-------------------------|-----------------------|-----------|-----------|
| Caucasian Proband (II-1) | 8 yr | Seize up Spasticity Progressive dystonia | Normal | Normal | Bilateral lesions in the putamen | Heteroplasmic | Pediatric onset dystonia (6) |
| II-2 | 19 yr | Cecocentral scotoma Asymptomatic | Present | Normal | Not tested | Heteroplasmic | LHON Asymptomatic |
| II-3 | 56 yr | | | | | |
| Caucasian Proband | 16 yr | Deteriorated visual activity | | Normal | Present | Normal | Heteroplasmic | LHON (6) |
| Japanese Proband (III-3) | 4 yr | Progressive dystonia, short stature Gradually deteriorated | Present | Bilateral lesions in the putamen | Heteroplasmic | G14459A combined of A3203G | LHON plus dystonia (7) |
| III-5 | 4 yr | Progressive dystonia, short stature Gradually deteriorated | Present | 17 yr | Not tested | Heteroplasmic | G14459A combined of A3203G | LHON plus dystonia |
| Korean Proband (proband), (II-12), (II-13), (III-14), (IV-3) | About 5 yr | Progressive dystonia | Delayed | Normal | Bilateral lesions in the putamen and caudate nucleus | Homoplasmic | Pediatric onset dystonia This report |
| II-7 | 55 yr | Asymptomatic | Normal | Normal | Not tested | Homoplasmic | Asymptomatic |
| III-11 | 36 yr | | | |

LHON, Leber hereditary optic neuropathy.

Table 2. The summary of clinical features of reported cases in Table 1 according to the described heteroplasmic or homoplasmic G14459A mutation status

| Parameters | Heteroplasmic mutation | Homoplasmic mutation | P value |
|------------|------------------------|----------------------|---------|
| Number of cases | 9 (34.4%) | 17 (65.4%) | |
| Mean age at onset | 17.8 yr (3-56) | 16.1 yr (2-55) | NS |
| Dystonia | 5 (55.6%) | 10 (58.8%) | NS |
| Impaired intelligence | 3 (37.5%) | 7 (46.7%) | NS |
| Ophthalmopathy | 6 (66.7%) | 3 (17.6%) | 0.012 |
| Basal ganglia degeneration of neuro image | 4 (44.4%) | 12 (70.6%) | NS |
| Clinical phenotype | Pediatric-onset dystonia | LHON | LHON plus dystonia | Asymptomatic |
| P value | 2 (22.2%) | 10 (58.8%) | 3 (33.3%) | 3 (17.6%) | 1 (11.1%) | 4 (23.5%) |

The statistical data were obtained using an SPSS software package, version 11.5 (SPSS, Chicago, IL, USA). A cut-off P value of 0.05 was adopted for all the statistical analyses. LHON, Leber hereditary optic neuropathy; NS, statistically non-significant.
found in three unrelated families with Leigh disease (13). This report adds to the clinical heterogeneity associated with the G14459A mtDNA mutation. The heterogeneous clinical features among the family members described here are unlikely to have arisen from different secondary mtDNA mutations among the family members because they were all from the same maternal lineage. Therefore, these findings support that other nuclear modifier genes are involved in the pathogenesis of the G14459A mutation (5).

In summary, the G14459A mutation is a candidate mutation for maternally inherited dystonia with bilateral striatal necrosis, regardless of optic neuropathy. To our knowledge, this is the first case revealing a mtDNA mutation in a Korean family with a maternally inherited dystonia. Moreover, the clinical manifestations of the homoplasmic G14459A mtDNA mutation, even within the same family, are heterogeneous and support the hypothesis that nuclear genes may play a role in modifying the clinical expression of mitochondrial disease.

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