Pediatric tetrasomy 18p presenting as a spastic cerebral palsy: A case report

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Tetrasomy 18p is a genetic syndrome caused by an isochromosome consisting of two copies of the short arm of chromosome 18. Clinically, pediatric cases of tetrasomy 18p manifest with global developmental delay, similar to most cases of chromosomal abnormality. In addition, it causes various symptoms including abnormal muscle tone. We report a case of an infant with global developmental delay and remarkable spasticity, the typical phenotype of bilateral spastic cerebral palsy. However, she had a subtle anomaly in her face, and brain magnetic resonance imaging (MRI) findings were inconsistent with her strong upper motor neuron signs. Upon genetic testing, she was determined to have an 18p isochromosome, confirming de novo non-mosaic tetrasomy 18p. Cerebral palsy is a neurological disorder that includes developmental delay caused by a non-progressive lesion in the developing brain. During diagnostic workup in patients with cerebral palsy, genetic testing should be considered when there are minor physical anomalies or equivocal MRI findings.

Key words: Tetrasomy 18p, Cerebral palsy, Hereditary spastic paraplegia.

Introduction

Tetrasomy 18p or isochromosome 18p (i[18p]) syndrome (Online Mendelian Inheritance in Man #614290) is caused by an isochromosome, a supernumerary marker chromosome, consisting of two copies of the short arm of chromosome 18. Even though 18p is the most common location of isochromosomes [1], it is still a very rare condition with a prevalence of less than 1 in 40,000 [2]. This syndrome affects both sexes equally, and most are de novo cases [3]. According to a literature review, global developmental delay has been observed in all reported 18p cases. Other features include abnormal muscle tone, seizures, and neonatal feeding problems, which are also frequent symptoms of cerebral palsy [3,4].

Recently, we encountered a pediatric case of de novo non-mosaic tetrasomy 18p, clinically diagnosed as spastic cerebral palsy by acquired brain lesion at another hospital. According to a literature review up to 2020, only one such case has been found in Korea, and since this subject had been prenatally diagnosed, their clinical manifestations were not reported [5].
Case

An 8-month-old female visited our department for physical therapy due to delayed motor development. As she was not able to creep forward and had severe spasticity affecting both lower extremities, she was clinically diagnosed with bilateral spastic type of cerebral palsy in another hospital. And acquired brain injury was suspected as the cause of cerebral palsy.

She was born by normal delivery at 41 weeks of gestation with a weight of 2.6 kg (below 3rd percentile) [6] without any history of trauma or hypoxic events. She was the second child of healthy parents, after her typically developing older sister. At the time of birth, her Korean father was 24 years old, and her Malian mother was 26 years old.

At 8 months of age, her body weight was 7.2 kg (10th–25th percentile), height was 72.6 cm (90th–97th percentile), and head circumference was 41 cm (below 5th percentile, microcephaly). Facial appearance showed low set ear and a smooth philtrum (Fig. 1), which were not found in other family members. Opisthotonic posture and spasticity were observed bilaterally, and we graded her hip adductors as grade 2, hamstrings as grade 1+, and heel cords as grade 2 as per the modified Ashworth Scale (MAS). On Bayley Scales of Infant and Toddler Development–2nd edition (BSID-II) evaluation at 9 months [7], her cognitive and psychomotor developmental scores were equivalent to those at 5 months and 4 months of age, respectively.

Brain magnetic resonance imaging (MRI) was performed at 9 months to rule out the possibility of brain lesions, and volume loss of the periventricular white matter and corpus callosum in the parietal area was suspected. Although it mimicked periventricular leukomalacia, the severity of the MRI could not fully explain with clinical severity (Fig. 2).

Electroencephalography showed a few bursts of semirhythmic delta activity on the right side of the head, suggesting focal cerebral dysfunction. Flash visual evoked potential showed delayed latency of P100 waves on both sides. Similar to the gastrointestinal problems in tetrasomy 18p which are reported previously [3,4], she also had severe constipation requiring intermittent lactulose syrup administration. Echocardiography and electrocardiogram screening revealed grossly normal cardiac anatomy with left ventricular hypertrophy. The clinical features of tetrasomy 18p reported in the literature in comparison with the findings in this case are summarized in Table 1.

Fig. 1. Dysmorphism observed in the patient. (A) Smooth philtrum. (B) Low set ear.

Fig. 2. Brain MRI (T1-weighted, axial [A] and sagittal [B]). Suspected decreased white matter volume in parietal areas, with a small splenium and posterior body of corpus callosum.
1. Genetic study

Genetic study was performed to identify the genetic cause of the dysmorphic features. As the first step, we performed high-resolution chromosome study and multiplex ligation-dependent probe amplification (MLPA) using a subtelomeric probe set (SALSA P070; MRC Holland, Amsterdam, Netherlands) [8], in which a small supernumerary marker chromosome was observed in every metaphase cell. The MLPA results showed a peak ratio of 2.0 at the 18p11.32 region, which strongly suggested that the marker chromosome originated from the short arm of chromosome 18, suggesting tetrasomy 18p (+i[18p]). Fluorescence in situ hybridization (FISH) analysis was conducted using a CEP 18 (D18Z1) SpectrumOrange probe (Abbott Laboratories, Des Plaines, IL, USA) to confirm the diagnosis. On FISH analysis, the supernumerary monocentric marker chromosome showed positive signals (Fig. 3). Consistent with most 18p isochromosome cases being de novo cases, the chromosomal analyses of the parents showed normal karyotypes.

2. Clinical progress with rehabilitation therapy

The study patient has received rehabilitation therapy at the

Table 1. Comparison of our case with the known phenotype of tetrasomy 18p

| Clinical finding               | Literature\(^a\) | This case | Total |
|-------------------------------|-----------------|----------|-------|
| Developmental delay           | 108/108         | Yes      | 109/109 |
| Dysmorphic features           | 108/108         | Yes      | 109/109 |
| Smooth philtrum              | 27/31           | Yes      | 28/32  |
| Small ears                    | 16/31           | No       | 16/32  |
| Abnormal muscle tone (hypertonia or hypotonia) | 79/108 | Yes | 80/109 |
| Congenital heart disease      | 23/98           | No       | 23/99  |
| Abnormal brain MRI            | 11/17           | Yes      | 12/18  |
| Myelomeningocele              | 6/108           | No       | 6/109  |
| Seizure                       | 23/108          | No       | 23/109 |
| Microcephaly                  | 51/108          | Yes      | 52/109 |
| Strabismus                    | 41/90           | No       | 41/91  |
| Recurrent otitis media        | 38/108          | No       | 38/109 |
| Constipation                  | 35/108          | Yes      | 36/109 |
| Neonatal complication         | 70/108          | No       | 70/109 |
| Growth retardation            | 30/108          | No       | 30/109 |
| Hearing loss                  | 11/94           | No       | 11/95  |

MRI, magnetic resonance imaging.

\(^a\)Values are presented as data from references [3]/[4].

Data from the article of Sebold C et al. (Am J Med Genet 2010;152:2164-72) [3]; Bawazeer S et al. (Appl Clin Genet 2018;11:9-14) [4].

Fig. 3. Genetic testing results. (A) High-resolution chromosome study showed an additional small metacentric marker chromosome. (B) Multiplex ligation-dependent probe amplification results (SALSA P070 subtelomeric probemix) indicated duplication of the 18p11.32 region. (C) Fluorescence in situ hybridization using CEP 18 (D18Z1) SpectrumOrange probe showed that monocentric marker chromosome was of chromosome 18 origin. Results summary: 47, XX, +mar.ish i(18)(p10)(D18Z1+).
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Since her hypertonus was considered the biggest problem, we tried antispastic medications. Dantrolene sodium was orally administered at 2 mg/kg daily from 10 months of age. It was changed to baclofen after 9 months because of non-significant efficacy of dantrolene sodium. Thus, she was orally administered 5 mg of baclofen three times per day for 19 months, when her heel cord spasticity increased to MAS grade 3, with equinus progression. Due to sufficient amelioration of spasticity, periodic motor point block at both gastrocnemius muscles using botulinum toxin A (3 IU/body weight in kg of onabotulinum toxin A [Botox®, Allergan, Allergan Sales LLC {Bioscience Laboratories} Campbell, CA, USA] in each muscle) has been performed since 31 months of age. Spasticity of both heel cords decreased from grade 3 to grade 2 after the first botulinum toxin injection. Additionally, we applied bilateral hinged ankle foot orthoses from 19 months of age to correct equinus gait patterns and facilitate walking. In order to improve gait ability, physical therapy that purposes to enhance balance and lower extremity uses with strengthening exercise has conducted. Occupational therapy purposed to facilitate eye-hand coordination and to stimulate visual perception including puzzle assembly and shape-matching has also provided.

3. Ethics statement
The study was approved by the Institutional Review Board of CHA Bundang Medical Center (2020-06-011). The images were published in agreement with the patient’s parents.

Discussion
To our knowledge, this is the first case of tetrasomy 18p in Korea, diagnosed during the workup of developmental delay and showing clinical progress. Previously reported case was discovered during prenatal workup for cardiac defects; thus, clinical courses thereafter was not reported, except for swallowing difficulty [5].

In previous studies, about 63% of patients showed abnormal brain MRI findings, including thin corpus callosum, brain atrophy, and lateral ventricle enlargement [3]. The study patient had a small periventricular, white matter which has not been reported previously. Although the various neurological features of tetrasomy 18p are seen, brain MRI findings are not frequently reported.

In the present case, the patient’s clinical symptoms were very similar to those seen in typical spastic bilateral cerebral palsy caused by periventricular leukomalacia, so she had been undergoing physical therapy in another hospital without further diagnostic workup. However, since the perinatal history and MRI findings could not completely explain the relatively severe clinical features, we conducted a gene study and were able to make...
the appropriate diagnosis. 

At the initial stage of diagnostic workup, a genetic condition called hereditary spastic paraplegia (HSP) was considered for diagnosis. HSP is known to be caused by over 80 genetic variants and is often diagnosed through new generation sequencing panel tests [9]. It is not easy to conduct tests in all suspected cases because of high cost. This report suggests that high-resolution chromosomal testing and MLPA may be helpful in some cases presenting with similar symptoms. All isochromosome 18p cases identified to date have been monocentric; the majority are de novo cases, and a few familial cases of maternal and paternal inheritance have been reported [10,11]. There are several mechanisms by which chromosome abnormality occur, and meiosis II nondisjunction, followed by subsequent postzygotic misdivision are the most common mechanisms [10]. In this study, no additional tests such as DNA polymorphism were performed, and the mechanism of isochromosome formation could not be elucidated. Only normal karyotype of the parents revealed that she was a de novo case.

In conclusion, this case implies that we should not overlook the clinical presentation of hypertonicity and be aware of the possibility of genetic causes in pediatric patients with developmental delay presenting with the phenotype of cerebral palsy. Subtle physical anomalies and brain image results may also be important clues for diagnosis.

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Authors’ Contributions

Conception and design: MYK. Acquisition of data: HK, WP, IL. Analysis and interpretation of data: IL, SHP. Drafting the article: IL, SHP. Critical revision of the article: SHS, MRS, MYK. Final approval of the version to be published: all authors.

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