Niemann-Pick Disease Type C Misdiagnosed as Cerebral Palsy: A Case Report

Eun Jae Ko, MD, PhD¹, In Young Sung, MD, PhD², Han-Wook Yoo, MD, PhD³

¹Department of Physical Medicine and Rehabilitation, Ulsan University Hospital, University of Ulsan College of Medicine, Ulsan; ²Department of Rehabilitation Medicine, Asan Medical Center, University of Ulsan College of Medicine, Seoul; ³Department of Pediatrics, Asan Medical Center Children’s Hospital, University of Ulsan College of Medicine, Seoul, Korea

Niemann-Pick disease type C (NP-C) is a rare autosomal recessive neurovisceral lysosomal lipid storage disorder. The clinical manifestations of the disorder are variable. This report describes the case of a 27-month-old girl with NP-C whose condition had been misdiagnosed as spastic cerebral palsy (CP). She had spasticity, particularly at both ankles, and gait disturbance. Magnetic resonance imaging of the brain revealed findings suspicious of sequelae from a previous insult, such as periventricular leukomalacia, leading to the diagnosis of CP. However, she had a history of hepatosplenomegaly when she was a fetus and her motor development had deteriorated, with symptoms of vertical supranuclear gaze palsy, cataplexy, and ataxia developing gradually. Therefore, NP-C was considered and confirmed with a genetic study, which showed mutation of the NPC1 gene. Thus, if a child with CP-like symptoms presents with a deteriorating course and NP-C-specific symptoms, NP-C should be cautiously considered.

Keywords Cerebral palsy, Niemann-Pick disease type C, Muscle spasticity
of a patient with NP-C presenting with gait disturbance with spasticity at both ankles, whose condition was initially misdiagnosed as spastic cerebral palsy (CP).

CASE REPORT

A 27-month-old girl visited the outpatient clinic of the Department of Pediatric Orthopedic Surgery for gait disturbance beginning at the age of 26 months. She showed mild limping with an intoeing gait pattern and spasticity at both ankles. Brain magnetic resonance imaging (MRI) at the age of 29 months revealed findings suspicious of sequelae from a previous insult, such as periventricular leukomalacia (Fig. 1), leading to the diagnosis of CP.

The patient was born at the gestational age of 37+3 weeks, with a weight of 2,035 g (<3%tile), height of 44 cm (3%tile–10%tile), head circumference of 31.5 cm (3%tile–10%tile), and Apgar score of 8>9, through spontaneous delivery. She was the first child, and her family history was not remarkable. Fetal ultrasonography revealed splenomegaly, hepatic congestion, echogenic bowel, and fetal ascites. The ascites resolved after abdomino-amniotic shunt operation in the fetal stage. After birth, she was treated for intrahepatic cholestasis. Her hepatosplenomegaly was continuously observed with ultrasonography. Tandem mass spectrometry screening and amino acid analysis showed nonspecific findings at 1 month.

She visited the outpatient clinic of the Department of Pediatric Rehabilitation Medicine and was admitted at the age of 31 months. Physical examination revealed spasticity in the lower extremities, particularly at the ankles (right<left), which was believed to be a symptom related to mild CP. She had global developmental delay (Table 1) and underwent rehabilitation treatment. Botulinum toxin injection targeting the left equinovarus foot and use of an ankle-foot orthosis helped with her gait pattern.

Neurological symptom deterioration was discovered incidentally via a video of the patient walking before the age of 2 years, which showed better gait pattern than that at the age of 31 months. Therefore, progressive diseases other than CP were considered. However, tests for arylsulfatase A enzyme for metachromatic leukodystrophy, very long-chain fatty acids for adrenoleukodystrophy, and beta-galactosylceramidase for Krabbe disease showed normal findings.

A few weeks later, the patient experienced sudden collapse of the knees when walking, which was first interpreted as an epileptic crisis; however, there was no abnormal finding on electroencephalography. The frequency of knee-collapse events increased, and cataplexy was gradually suspected because knee collapse also occurred when laughing. Furthermore, vertical supranuclear gaze palsy (VSGP) and wide ataxic gait pattern appeared progressively. The NPC1 gene for NP-C showed a mutation in a genetic study, and NP-C was diagnosed.

Table 1. Results of Denver Developmental Screening Test

|                      | At 31 months | At 47 months |
|----------------------|--------------|--------------|
| Personal-social (mo) | 18–19        | 6–7          |
| Fine motor-adaptive (mo) | 8–9        | 7–8          |
| Receptive language (mo) | 18–20     | 6–7          |
| Expressive language (mo) | 10–11     | 6–7          |
| Gross motor (mo)     | 12–13        | 6–7          |

Fig. 1. Brain magnetic resonance image showing (A) mild bilateral T2/fluid-attenuated inversion recovery hyperintensities in the peritrigonal periventricular white matter and (B) mild thinning of the corpus callosum with features suspicious of mild hypoplastic splenium.
From the age of 34 months, treatment for NP-C, including oral miglustat and intrathecal and intravenous betacyciclodextrin, was initiated and continued. However, her clinical manifestations continued to worsen (Table 1). The informed consent was waived for this case report.

**DISCUSSION**

Differentiating slowly progressive neurological disorders from CP can be challenging, because these conditions frequently share similar patterns of motor impairment. CP describes a group of permanent disorders of movement and posture that are attributed to non-progressive disturbance of the developing fetal or infant brain [4]. The key characteristic distinguishing NP-C from CP is the progressive pattern of the disease. Furthermore, some uncommon features of NP-C may help in diagnosing this condition, including visceral, neurological, and psychiatric symptoms.

Visceral symptoms are the most specific symptoms of NP-C [5]. Splenomegaly is the strongest visceral indicator of the disease [1]. Hepatomegaly is observed less frequently than splenomegaly. Prolonged or unexplained neonatal cholestatic jaundice is also a strong visceral indicator of NP-C [5]. The presence of fetal ascites or hydrops fetalis is less frequent [5].

Neurological symptoms are the most common and sensitive indicators of NP-C [5]. VSGP occurs owing to selective vulnerability of neurons at the rostral interstitial nucleus of the medial longitudinal fasciculus and causes deficits in voluntary and reflexive vertical saccades as well as vestibulo-ocular nystagmus [6]. Cataplexy is a sudden episode of muscle weakness with full conscious awareness, triggered by laughing, crying, or the emotion of fear, which can appear at as early as 2 years of age [7]. It is frequently misinterpreted as a fall or as atonic epileptic crises [1]. VSGP and cataplexy are the strongest neurological indicators of NP-C [5]. Slow ataxia is a moderate indicator of NP-C and becomes highly suggestive of NP-C in combination with dystonia of the hands and face [5]. Hypotonia is usually observed in the second year, and spasticity is sometimes observed in the early or late infantile periods. Developmental delay, dysarthria, and dysphagia can also occur [5].

Psychiatric symptoms are the least specific indicators of NP-C [5]. All patients present with progressive cognitive decline, which is a strong indicator of the disease [5]. Psychotic symptoms (hallucination or delusion) are moderate indicators of NP-C [5] and typically appear during adolescence or early adulthood.

The radiographic features of NP-C are nonspecific, among which cerebral (particularly frontal lobe) and cerebellar atrophy [3,8], high T2-weighted signal intensity in the parieto-occipital periventricular white matter [8], deep gray matter and hippocampal atrophy in adult-onset NP-C [9], and reduced midbrain-to-pons ratio [10] have been reported. However, the patient with NP-C in this report did not show the above-mentioned findings on brain MRI.

Among the above features of NP-C, the patient in this case report showed hepatosplenomegaly, cataplexy, VSGP, and spasticity; however, interestingly, she did not show symptoms of dystonia, hypotonia, cognitive decline, hallucination, and delusion. Therefore, this case is different from typical NP-C cases.

In conclusion, this report highlights the importance of suspecting other diseases when deterioration of symptoms is detected in a patient with CP-like symptoms, because early treatments for metabolic diseases are important to stabilize or slow disease progression. Therefore, when a patient with CP-like symptoms presents with a deteriorating course and NP-C-specific symptoms, NP-C should be cautiously considered.

**CONFLICT OF INTEREST**

No potential conflict of interest relevant to this article was reported.

**AUTHOR CONTRIBUTION**

Conceptualization: Ko EJ, Sung IY, Yoo H. Methodology: Ko EJ, Sung IY. Formal analysis: Ko EJ, Sung IY, Yoo H. Project administration: Sung IY. Visualization: Ko EJ. Writing -original draft: Ko EJ. Writing - review and editing: Sung IY. Approval of final manuscript: all authors.

**REFERENCES**

1. Patterson MC, Hendriksz CJ, Walterfang M, Sedel F, Vanier MT, Wijburg F, et al. Recommendations for the diagnosis and management of Niemann-Pick disease
type C: an update. Mol Genet Metab 2012;106:330-44.
2. Lloyd-Evans E, Platt FM. Lipids on trial: the search for the offending metabolite in Niemann-Pick type C disease. Traffic 2010;11:419-28.
3. Vanier MT. Niemann-Pick disease type C. Orphanet J Rare Dis 2010;5:16.
4. Rosenbaum P, Paneth N, Leviton A, Goldstein M, Bax M, Damiano D, et al. A report: the definition and classification of cerebral palsy April 2006. Dev Med Child Neurol Suppl 2007;109:8-14.
5. Wijburg FA, Sedel F, Pineda M, Hendriksz CJ, Fahey M, Walterfang M, et al. Development of a suspicion index to aid diagnosis of Niemann-Pick disease type C. Neurology 2012;78:1560-7.
6. Salsano E, Umeh C, Rufa A, Pareyson D, Zee DS. Vertical supranuclear gaze palsy in Niemann-Pick type C disease. Neurol Sci 2012;33:1225-32.
7. Nevsimalova S, Malinova V. Cataplexy and sleep disorders in Niemann-Pick type C disease. Curr Neurol Neurosci Rep 2015;15:522.
8. Huang JY, Peng SF, Yang CC, Yen KY, Tzen KY, Yen RF. Neuroimaging findings in a brain with Niemann-Pick type C disease. J Formos Med Assoc 2011;110:537-42.
9. Walterfang M, Patenaude B, Abel LA, Kluenemann H, Bowman EA, Fahey MC, et al. Subcortical volumetric reductions in adult Niemann-Pick disease type C: a cross-sectional study. AJNR Am J Neuroradiol 2013;34:1334-40.
10. Walterfang M, Macfarlane MD, Looi JC, Abel L, Bowman E, Fahey MC, et al. Pontine-to-midbrain ratio indexes ocular-motor function and illness stage in adult Niemann-Pick disease type C. Eur J Neurol 2012;19:462-7.