Niemann-Pick disease type C: a case series of Brazilian patients

Doença de Niemann-Pick tipo C: série de casos de pacientes brasileiros

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ABSTRACT

The aim of the study was to analyze a series of Brazilian patients with Niemann-Pick disease type C (NP-C). Method: Correlations between clinical findings, laboratory data, molecular findings and treatment response are presented. Result: The sample consisted of 5 patients aged 8 to 26 years. Vertical supranuclear gaze palsy, cerebellar ataxia, dementia, dystonia and dysarthria were present in all cases. Filipin staining showed the "classical" pattern in two patients and a "variant" pattern in three patients. Molecular analysis found mutations in the NPC1 gene in all alleles. Miglustat treatment was administered to 4 patients. Conclusion: Although filipin staining should be used to confirm the diagnosis, bone marrow sea-blue histiocytes often help to diagnosis of NP-C. The p.P1007A mutation seems to be correlated with the "variant" pattern in filipin staining. Miglustat treatment response seems to be correlated with the age at disease onset and disability scale score at diagnosis.

Keywords: Niemann-Pick disease type C, bone marrow, filipin stain, NPC1 gene, miglustat.

RESUMO

O objetivo desse estudo foi analisar uma série de casos de pacientes brasileiros com doença de Niemann-Pick tipo C (NP-C). Método: Correlação entre manifestações clínicas, alterações laboratoriais, estudo molecular e resposta ao tratamento foram realizadas. Resultado: A amostra consiste de 5 pacientes com idade entre 8 e 26 anos. Paralisia do olhar vertical supranuclear, ataxia cerebelar, demência, distonia e disartria estavam presentes em todos os casos. Coloração de filipina na cultura de fibroblastos mostrou padrão “clássico” em dois pacientes e padrão “variante” em três casos. O estudo molecular encontrou mutações no gene NPC1 em todos os alelos. O tratamento com miglustate foi realizado em 4 pacientes. Conclusão: Embora coloração de filipina seja utilizada para confirmar o diagnóstico, o histiócito azul-marinho no aspirado de medula óssea frequentemente auxilia a confirmar o diagnóstico de NP-C. A mutação p.P1007A está correlacionada com o padrão “variante” na coloração de filipina. A resposta ao tratamento com miglustate parece estar correlacionada com a idade e escorre de desabilidade no momento do diagnóstico.

Palavras-chave: doença de Niemann-Pick tipo C, medula óssea, coloração de filipina, gene NPC1, miglustate.

Niemann-Pick disease type C (NP-C) is a rare inherited disease, caused by mutations in either the NPC1 or the NPC2 gene, which leads to impaired intracellular lipid trafficking and the accumulation of cholesterol and glycosphingolipids in the brain and other tissues¹-⁶. The clinical signs and symptoms of NP-C can develop at any age, and significant phenotypic heterogeneity is frequently observed in NP-C¹²-⁴⁷. Vertical supranuclear gaze palsy, cerebellar ataxia, dystonia, dementia, epilepsy and visceral manifestations are the most common symptoms of NP-C¹²-⁴⁷.

NP-C patients often show sea-blue histiocytes or foamy cells in bone marrow analyses and intracellular lipid accumulation with filipin staining¹,³,⁴,⁶. Molecular analyses typically show NPC1 (in 95% of cases) or NPC2 (in around 4% of cases) gene mutations¹⁴,⁴⁶. Nevertheless, vertical supranuclear gaze palsy, cataplexia, high chitotriosidase serum
level and relatives with similar symptoms are also considered to be clinical clues pointing to the diagnosis4-7.

In this study, we analyzed clinical manifestations, disease progression, brain imaging, laboratory features, intracellular lipid accumulation and molecular findings from 5 patients with NP-C to contribute to the better characterization of this disease in Brazil.

METHOD

A retrospective analysis of five patients with NP-C, diagnosed between January 2005 and January 2013, was performed based on the following features: typical clinical manifestations; foamy cells or sea-blue histiocytes cell in the bone marrow; or abnormal intracellular lipid accumulation observed with filipin staining. Informed consent for laboratory and molecular analyses were obtained from all of the patients (in the outpatient clinic or during hospital admission for diagnostic investigation in the Hospital de Clínicas da Universidade Federal do Paraná).

Clinical evaluation

Relevant data were collected, including age, gender, neurological and visceral manifestations, clinical evaluation, and the presence of other affected relatives. The clinical presentation was categorized according to the patient's age at onset of the disease as follows: neonatal (onset at age ≤3 months), early infantile (onset at age 3 months to 2 years), late infantile (onset at age 2 to <6 years), juvenile (onset at age 6 to 15 years) or adult (onset at age >15 years). The NP-C-specific disability scale was used to evaluate the severity of the disease3. Patient response to substrate reduction therapy with miglustat was classified as present, absent, or partial based upon the objective improvements in clinical manifestations and the scores on the NP-C-specific disability scale over the course of the drug treatment (after at least 1 year of use).

Brain images

Radiologic findings from brain magnetic resonance imaging (MRI) were also reported.

Laboratory analysis

Chitotriosidase levels in patient serum samples were measured (normal range: 8.8 – 132.0 nmol/h/ml).

Bone marrow findings

Bone marrow aspiration and biopsy specimens were analyzed to identify sea-blue histiocytes (Niemann-Pick cells) with May-Grünwald-Giemsa staining and foamy cells with hematoxilin-eosin staining.

Filipin test

Skin biopsy samples were used for fibroblast cultures, and cells were stained for histological examination. The presence of intracellular lipid accumulation was identified with filipin staining; in the ‘classical’ cholesterol storage pattern, positive cells are typically strongly fluorescent (cholesterol-filled perinuclear vesicles). The pattern of the cell samples was categorized with filipin staining as either normal (clear negative fluorescence); atypical or “variant” (moderated fluorescence); or typical or “classical” (high fluorescence).

Molecular analysis

Blood samples from the patients were used to isolate DNA by standard method. Coding sequences and flanking regions of the NPC1 and the NPC2 genes were amplified with PCR, purified and submitted to direct DNA sequencing using the BigDye™ Terminator Cycle Sequencing kit v. 3.1 (Applied Biosystems, Foster City, CA, USA) following the manufacturer’s instructions. Products were then submitted to capillary electrophoresis in an ABI PRISMTM 3130xl Genetic Analyzer, and sequences were analyzed with DNA Sequencing Analysis software v. 5.2 (Applied Biosystems). Mutations were confirmed by sequencing an independent DNA sample with both forward and reverse primers.

RESULTS

The report includes five patients (four females and one male) aged 8 to 26 years. The time of disease progression ranged from 2 to 16 years, with a mean time of 6 years. Family members with the same disease were not found. The age at onset of the disease was early infantile in one patient, late infantile in one patient and juvenile in three patients (Table).

All patients had dementia, dystonia, cerebellar ataxia, dysarthria and vertical supranuclear gaze palsy. Other reported clinical features included the following: psychiatric disorders (4/5), epilepsy (2/5), dysphagia (4/5), dysphonia (4/5), weakness (4/5), hepatosplenomegaly (3/5), cataplexy (2/5), cholestasis (1/5) and deafness (1/5) (Table).

Brain MRI was performed for all patients (Table). Diffuse cerebral and cerebellar atrophy was observed in three patients, while two patients had normal imaging results. Mild multifocal areas of hypersignal in the cerebral white matter were observed in one patient (Case 1).

Serum chitotriosidase activity was measured in 4 patients (Table). Serum chitotriosidase levels were above normal limits in all patients and varied from 183 to 1477 nmol/h/ml (mean level of 547.75 nmol/h/ml).
Bone marrow specimens were analyzed in 4 patients (Table). Sea-blue histiocytes and foamy cells were identified in three patients (Figures D, E and F).

Skin fibroblast culture was performed for all patients (Table). The cell patterns from the filipin stains were classified as atypical in three patients and typical in two patients (Figures A, B and C).

Molecular analysis was performed in all patients (Table). Mutations in \textit{NPC1} gene were found in all alleles. Two patients were homozygotes for the p.P1007A mutation, and compound heterozygotic mutations (p.P1007A, p.A1035V, p.A764V, p.S954L and p.P887L) were found in three patients.

Substrate reduction therapy with miglustat (600 mg/day) was administered to 4 patients (Table). The treatment response was clearly beneficial in at least one case (Case 4). The miglustat dosage was decreased (to 200 mg/day) due to the occurrence of gastrointestinal side effects in one patient (Case 2).

Disease follow-up after diagnosis ranged from 2 to 16 years with a mean time of 6.6 years. NP-C-specific disability scale scores ranged from 7 to 15 (mean: 10.2) at disease diagnosis and from 7 to 18 (mean: 12.6) at final follow-up. One patient died during the treatment due to pneumonia (Case 3).

**DISCUSSION**

Although specific clinical features vary, some main core features were present in all of our patients. Vertical supranuclear gaze palsy has been included as a clinical criterion.
or core feature of NP-C. This criterion may be an important clinical feature, which can help to identify the disease in patients with cerebellar ataxia, dystonia or dementia.

Brain MRI generally reveals cerebellar atrophy that can be associated with cerebral atrophy or unspecific abnormalities of cerebral white-matter. Brain MRI is important for initial screening and differential diagnoses of other causes of cerebellar ataxia, epilepsy and dementia, but the findings are not specific for the NP-C diagnosis. In addition, high serum chitotriosidase levels were useful as an initial screening for our patients, but this enzyme is not a specific marker for NP-C. Other assessments are necessary for disease diagnosis.

Diagnostic investigation revealed abnormal cell patterns after filipin staining in all patients, but the typical pattern was found in only two patients. The presence of an atypical pattern after filipin staining is associated with specific mutations of the NPC1 gene. The three cases presenting an atypical pattern after filipin staining all presented with the p.P1007A mutation. Our findings help to confirm previous reports suggesting that the p.P1007A mutation can show atypical or "variant" findings with filipin staining. We believe that in these cases, bone marrow analysis can be useful for NP-C diagnosis because the presence of sea-blue histiocytes and foamy cells was similar for all of our cases. Although sea-blue histiocytes are not pathognomonic of NP-C, the typical pattern of filipin staining was found for two patients and was important for the case with a new mutation (p.A764V) of the NPC1 gene to confirm that this was a pathogenic mutation. We believe that fibroblast culture to filipin staining is a technically complex procedure. We suggest that, in the future, this procedure should be used only to confirm the pathogenic character of a new mutation after molecular analysis. Therefore, filipin staining of the bone marrow cells can be used to provide a rapid screening test for NP-C, but it is not considered to be a definitive diagnostic method.

Several mutations of the NPC1 gene have been described in NP-C patients. The p.P1007A mutation of the NPC1 gene is frequently reported to cause, mainly juvenile or adult forms, of the disease in several countries. In our study in Brazil, only a few patients with molecular analysis results available were described, and it is not possible to conclude what is the most common mutation in this population. The p.P1007A and p.A1035V mutations were previously reported in Brazil, and our study contributes to the idea that they are frequent NPC1 mutations in this country although only a few patients were analyzed.

Treatment response can be evaluated with the NP-C-specific disability scale during treatment. In our patients, a clear beneficial response was observed in one case. We should note that disability scores at diagnosis were already high in cases which did not show observed beneficial response, which could have contributed to our results.
miglustat is less likely to provide substantial therapeutic benefits if severe neurological impairment is already present at diagnosis. In these cases, decisions to start treatment should be on a case-by-case basis. Comparisons between disability scores at diagnosis and treatment response in our study agree with the classical clinical opinion that treatment response is better in patients with mild neurological disability. As most of our patients had early onset of the disease, this circumstance could have influenced our findings as well, as treatment response is correlated with the age at onset, with adult onset being less disabling than infantile onset. To make available the diagnostic toolkits that are needed to provide earlier diagnosis is important goal in Brazil to obtain better treatment results for NP-C patients.

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