An Irish Patient With A Rare Leukodystrophy Related To Lamin B1 Duplication

Abstract:
A Molloy, O Cotter, R van Spaendonk, E Sisterrman, B Sweeney
Department of Neurology, Cork University Hospital, Wilton, Cork

The hereditary leukodystrophies are rare disorders caused by molecular abnormalities leading to destruction of or failure of development of central white matter. For almost 30 years there has been increasing recognition of later onset Autosomal Dominant Leukodystrophy (ADLD). We report the first genetically confirmed case of lamin B1 duplication causing ADLD from Ireland.

Case Report
A 47 year old gentleman presented with a six year history of urinary dysfunction including hesitancy, frequency and nocturia. He described increasing fatigue and hearing loss more marked on the left side over the preceding 18 months. In his past history he had had a severe upper gastrointestinal bleed secondary to duodenal ulcer with a marked hypotensive episode peri-operatively. Of note in his family history, his mother had died of a progressive neurological illness at age 57. This had been labelled as a brain tumour but no confirmatory evidence had been provided. Of three male siblings, two had died of myocardial infarction in their forties. Another had a diagnosis of bipolar affective disorder.

On initial examination, there were no cognitive deficits noted. He had an ataxic gait, Romberg’s sign was positive. Cranial nerves were intact. Tone and power were normal on limb examination. Reflexes were 2+ throughout and plantar responses were extensor. There was past-pointing suggesting cerebellar dysfunction bilaterally. Blood results were within normal limits or negative including: vitamin B12, folate, thyroid function tests, cortisol levels, very long chain fatty acids, HRV, HTLV-1, Lymedisease antibodies, ACE, Genetic testing for CADASIL was negative. Cerebrospinal Fluid analysis revealed normal cell count, cytology, protein, glucose with negative oligoclonal bands. An audiogram revealed asymmetric bilateral sensorineural hearing loss. Visual evoked responses showed dispersed low amplitude responses, consistent with possible optic nerve axonal loss. Electroretinogram showed mild delay bilaterally but acceptable amplitudes were noted, implying possible optic nerve pathology.

MRI brain scan showed widespread white matter disease in centrum semioval and both cerebral and cerebellar peduncles without enhancement. Long tract involvement was confirmed in midbrain, pons and medulla was noted. Over the succeeding 6 years he continued to progress with symptoms including increasing fatigue, ataxia and falls, dysarthria, and ongoing urinary frequency and urgency. Progression of neurological signs was noted including ataxia, dysarthria, apraxia and bilateral lower extremity weakness. MRI showed deterioration with extensive signal change in the centrum semiovale, pyramidal tracts, midbrain, cerebellar peduncles, medial lemnisci and signal changes in the lower cervical cord. Ventrices were dilated in keeping with a degree of atrophy (Figures 1 and 2). Because of the clinical and MRI findings, LMNB1 (Lamin B1) testing was performed. A positive result for LMNB1 exon 1 t/m 11 duplication was obtained consistent with a diagnosis of ADLD. MLPA analysis was used to detect the duplication, using the P071 kit from MRC-Holland (more detail available at http://mlpa.com).

Figure 1: Involvement of cerebellar peduncles on FLAIR MRI scan.

Figure 2: Diffuse predominately fronto-parial white matter changes on T2W imaging in advanced stages.

Discussion
This patient had an 8-10 year history of progressive multi-system neurological dysfunction with MRI showing a marked diffuse leukodystrophy. There was a family history of a progressive early onset neurological disorder in his mother which now raises the possibility of her being affected by ADLD. The diagnosis carries significant implications for these patients sibling and children, as it has been highly penetrant in the families described so far. ADLD due to Lamin B1 duplication was first described in 1984 by Eldridge et al in an Irish-American kindred. The first four cases had motor impairment and ataxia as in our patient. Slow progression of symptoms to death after 20 years was typical. MRI characteristics in kindreds described include diffuse white matter changes in a symmetrical distribution bilaterally, most prominent in fronto-parial areas and cerebellar peduncles. Spinal cord involvement was noted, as has cerebral, cerebellar and medullary atrophy.

Radiations initially is characteristic, unlike many other leukodystrophies. In our case the imaging findings were typical. ADLD is the first human disease known to be caused by a mutation of the gene responsible for lamin B1, and is the only laminopathy caused by over expression. Excess folding and blebbing of the nuclear envelope has been shown to occur at a cellular level, and the extent of expression is crucial in regulation of oligodendrocyte maturation and myelin formation. Worldwide at this point, nine families have been described, including two of Irish-American origin. Brusso et al reported an Italian kindred with a form of ADLD without LMNB1 duplication, the autonomic features were not present at the outset, raising the possibility that other genetic causes for variant forms of ADLD exist. We are unaware of any connection between previous cases and our patient at this time.

Correspondence: A Molloy
Department of Neurology, St Vincents University Hospital, Elm Park, Dublin 4
Email: a.molloy@st-vincents.ie

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References