Late-onset and acute presentation of Brown-Vialetto-Van Laere syndrome in a Brazilian family

Sarah Camargos, MD, PhD, Rita Guerreiro, MS, PhD, Jose Bras, MS, PhD, and Luis Sergio Mageste, MD

Neurol Genet 2018;4:e215. doi:10.1212/NXG.0000000000000215

Riboﬂavin transporter deﬁciency (formerly known as Brown-Vialetto-Van Laere [BVVL] or Fazio-Londe syndrome) is a neurodegenerative disorder characterized by progressive bulbar palsy with sensorineural deafness or bulbar hereditary neuropathy. It is caused by mutations in the riboﬂavin transporter genes SLC52A2 (RFVT2) or SLC52A3 (RFVT3). It is a rare syndrome with approximately 70 cases reported worldwide, with molecular diagnoses of RFVT2 or RFVT3. We have previously described the ﬁrst Brazilian family with a clinical diagnosis of BVVL.

In this report, we extend the clinical spectrum associated with this family and describe a new mutation related to the metabolism of riboﬂavin.

Case report

The proband was a previously healthy woman aged 34 years, who presented with hearing and vision loss in the last 6 months (ﬁgure, A). She was disturbed by facial pain, numbness in the left hemiface, diﬃculty moving her tongue, dysphagia, weight loss, and bilateral foot drop.

Examination demonstrated bilateral optic atrophy, normal ocular movements, bilateral facial paresis, atrophic tongue, and ﬂaccid dysarthria. Reﬂexes were brisk except for ankle reﬂexes that were absent. Plantar responses were indifferent. All sensory modalities were normal. Strength was globally diminished with important distal impairment and foot drop. As dysphagia and dyspnea progressed, a feeding tube was placed and noninvasive ventilation support was initiated. At that time, she was quadriplegic and could not walk.

Electroneuromyography demonstrated cervical and acute lumbar denervation, with chronic neurogenic changes. Audiologic evaluation demonstrated neurosensorial loss.

The patient was the eldest sibling of a consanguineous marriage. She had 3 maternal aunts, also sisters from a consanguineous marriage, with a probable diagnosis of BVVL syndrome.

The patient was started on empiric treatment with riboﬂavin (1,800 mg per day), and within 6 months of therapy, she could walk with a cane; the feeding tube and noninvasive ventilation were withdrawn (ﬁgure, B–G). Electroneuromyography was performed after B2 treatment and demonstrated low CMAP amplitudes and persistence of recent denervation (ﬁgure e-1, http://links.lww.com/NXG/A17).
Methods

The genetic study of the patient and both parents was conducted after written informed consent was obtained. Whole-exome sequencing (WES) was performed in the index case, and whole-genome genotyping (WGG) was used to identify large tracts of homozygosity in the trio, given the apparent autosomal recessive pattern of inheritance of the disease in the family and the presence of several loops of consanguinity. The 3 samples were genotyped at the whole-genome level using Illumina HumanOmniExpress Bead chips, and results were analyzed with GenomeStudio.

Results

By analyzing the variability identified by WES in genes previously known to cause riboflavin transporter deficiency, we identified a novel homozygous insertion in SLC52A3 (ENST00000217254:SLC52A3:c.1232_1233insCTACGC TTCCCTCCGCCCAGTGCTCGTGGTGTTCTTCACGCCGTCCTCACG; p.Ser411_Tyr412inS TyrAlaSerLeuProAlaProGlnAlaSerTrpValLeuPheSerGly CysLeuSer). The mutation was confirmed to be present in homozygosity in the index and was found in heterozygosity in both parents using Sanger sequencing. In addition, WGG revealed a large (1.5 Mb) homozygous region encompassing the SLC52A3 locus (chromosome 20: 643,919–2,146,580 Mb) that was not present in either parent. Consequently, we tested the phenotypically affected aunt (II.10), and she presented the mutation in homozygosity (figure e-2, http://links.lww.com/NXG/A18). The mutation found is neither present in homozygosity in population databases of genetic variability (ExAC and gnomAD), nor have we identified it in our in-house sequencing data from healthy controls and other diseases (n > 6,000).

Discussion

Both SLC52A3 and SLC52A2 are riboflavin transporters. Riboflavin is a precursor of flavin mononucleotide and flavin adenine dinucleotide, which are important cofactors for energy metabolism. Since the description of these 2 genes related to riboflavin transporter deficiency, the perspective about this disease has changed significantly.2,4,6 However, the mechanisms by which the disrupted proteins lead to the disease are still to be fully understood.

Motor neuron disease, neurosensory deafness, optic neuropathy, and the involvement of cranial nerves are common in both genotypes.4,7 Nevertheless, some phenotypic differences have been described: RFVT2 mutations present predominantly upper limb weakness and sensory ataxia; such findings are not commonly seen in patients with RFVT3 mutations.7

Here, we report a Brazilian patient with late-onset and uncharacteristic acute and severe presentation, demonstrating some phenotypic heterogeneity within a family.5 The mutation, a homozygous insertion of 60 bp in SLC52A3, has not been previously described as the cause of riboflavin transporter deficiency. So far, response to riboflavin therapy was documented
in 11 patients harboring mutations in RFVT3. Of them, 9 patients demonstrated some response and 2 remained stable.3,6 Some authors argue that response tends to be better and more rapid when earlier treatment is started. Riboflavin dose reposition is unknown, and treatment, although generally efficient, is empirical. In addition, there is still no evidence to reassure that treatment would prevent the occurrence of symptoms indefinitely. Despite all this, clinicians might be aware of this potentially treatable condition and initiate riboflavin supplementation as soon as diagnosis is suspected.

Author contributions
Sarah Camargos: study concept, design, acquisition of data, and study supervision. Rita Guerreiro: study concept, design, and critical revision of the manuscript for important intellectual content. Jose Bras: analysis and interpretation of the data. Luis Sergio Mageste: analysis and interpretation of the data.

Acknowledgment
The authors acknowledge the patients who participated in this study.

Study funding
No targeted funding reported.

Disclosure
S. Camargos has served on the scientific advisory board of Teva Pharmaceuticals and has received travel funding/speaker honoraria from Roche, Teva Pharmaceuticals, and Aché Pharmaceuticals. R. Guerreiro serves on the editorial boards of Science Matters, the American Journal of Neurodegenerative Disease, and the Journal of Parkinson’s Disease and has received research support from Alzheimer’s Research UK and Alzheimer’s Society. J. Bras serves on the editorial board of the Journal of Parkinson’s Disease and has received research support from Alzheimer’s Society. L.S. Mageste reports no disclosures. Full disclosure form information provided by the authors is available with the full text of this article at Neurology.org/NG.

Received July 19, 2017. Accepted in final form November 17, 2017.

References
Late-onset and acute presentation of Brown-Vialetto-Van Laere syndrome in a Brazilian family
Sarah Camargos, Rita Guerreiro, Jose Bras, et al.
Neurol Genet 2018;4;
DOI 10.1212/NXG.0000000000000215
This information is current as of January 26, 2018