Danon Disease in an Asymptomatic Woman: A Five-Year Follow Up

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Introduction

Danon disease (DD)¹ is a rare, dominant X-linked disease caused by mutation of the LAMP2 gene, which encodes a lysosome-associated membrane glycoprotein, thereby affecting lysosomal deposition. DD is characterized by a classic triad of cardiomyopathy (featured by hypertrophic cardiomyopathy [HC]), skeletal myopathy, and cognitive changes. While female patients tend to have milder phenotypic manifestations, an isolated cardiac involvement, in addition to a later onset of symptoms, without the need for heart transplantation before the fourth decade of life, male patients commonly have the presentation of the classic triad of disease.²

The clues of the involvement of HC with pre-excitation and persistent increased troponin I in these individuals are related to the process of autophagy that contributes to cardiac remodeling.³ However, there is still no specific treatment for DD. The approach to cardiac manifestations includes implantable cardioverter defibrillator (ICD) and ablation to improve symptoms and decrease the risk of sudden death. In cases of advanced heart failure (HF), heart transplantation is an effective and safe measure. Studies for gene therapy are currently in progress.⁴,5

Keywords

Glycogen Storage Disease Type II/genetics; Cardiomyopathy Hypertrophic; Phenotype; Lysosomal-Associated Membrane Protein 2/genetics (DD Danon Disease).

Considering the small number of cases described in the literature about DD and the gap in knowledge for an earlier approach, we aimed to describe the case of a patient with incidental diagnosis of DD, presenting a mutation not previously described in the literature and its five-year follow-up.

Case description

A female 23-year-old Caucasian patient, only daughter of a no consanguineous couple, was incidentally diagnosed with HC at 18 years of age during the preoperative period of an orthopedic surgery and confirmed by cardiac resonance. The parents were asymptomatic, with normal echocardiogram.

At the age of 20, she was admitted to the emergency department with atypical chest pain, and no other findings at physical examination, and laboratory tests showed an increase in troponin I (6ng/dL). She underwent a new cardiac magnetic resonance imaging confirming the diagnosis of CH with delayed gadolinium enhancement (Figure 1). At the time, she was diagnosed with acute myocarditis.

Patient was referred to a specialized HF center, where a Doppler echocardiography was performed, confirming the findings of HC (Figure 2), and revealing slight obstruction of the outflow pathway, and a Wolff-Parkinson-White (WPW) preexcitation pattern. The patient had elevated and stable levels of troponin I (7.74ng/dL), and natriuretic peptide (BNP) levels of 401 pg/mL six months after admission for chest pain. The patient underwent genetic testing; a new mutation NP_002285.1:p.Asn242Thrfs*41 compatible with DD was identified, and referred as accessory pathway (VA) ablation due to the WPW-type preexcitation.
Figure 1 – Gadolinium-enhanced cardiac magnetic resonance imaging showing left ventricular hypertrophy with involvement of apical segments and mid-basal portion of the anterior and lateral walls and left ventricular diastolic dysfunction, suggestive of myocardial fibrosis observed in cases of hypertrophic cardiomyopathy. Left ventricular ejection fraction: 78%. Maximum end diastolic thickness of septum IV = 2.1 cm; lateral wall = 2.0 cm.
Figure 2 – Doppler echocardiogram showing alteration of left ventricular diastolic function and hypertrophic cardiomyopathy
After three months, the patient presented with a new WPW-compatible pre-excitation episode involving the same VA, and referred for another ablation. A follow-up Holter revealed an asymptomatic, non-sustained ventricular tachycardia (VT) (Figure 3), and an implantable cardioverter defibrillator (ICD) was indicated, based on the cardiac resonance imaging findings also.

Since then, the patient has been asymptomatic, undergoing cardiac rehabilitation and treatment with beta-blockers. There has been no ICD firing since its implantation, in addition to normal renal, hepatic, ophthalmic and neurological functional tests. The patient has been followed by the departments of clinical genetics, cardiology, and arrhythmology, and received psychological support for anxiety disorder. In the last months she has been in isolation due to the COVID-19 pandemic and routinely performed physical activities and in telemedicine consultation. The temporal progression of the events were described in Tablet 1.

![Figure 3 – Electrocardiogram showing non-sustained ventricular tachycardia.](image)

| Temporal Evolution | Events |
|--------------------|--------|
| 2014 18 years old  | Diagnosis of hypertrophic cardiomyopathy during the preoperative evaluation for orthopedic surgery. |
| 2016 20 years old  | Patient admitted for atypical chest pain. The patient was referred to a specialized cardiac failure service that identified in 12-lead ECG the presence of Wolff-Parkinson-White preexcitation pattern. |
| 2017 21 years old  | Patient seeks the emergency for the second time with complaint of palpitation; she was diagnosed with Wolff-Parkinson-White pre-excitation and referred to first ablation. |
| 2018 22 years old  | After the insertion of the Holter track, several episodes of non-sustained ventricular tachycardia (VT) were observed. The patient was referred for implantable cardioverter defibrillator placement. |
| 2019 23 years old  | The patient remains asymptomatic, undergoing multidisciplinary follow-up and cardiac rehabilitation three times a week. In the last months she has been in isolation due to the COVID-19 pandemic and routinely performing physical activities and in telemedicine consultation. |
Discussion

The present study reports the case of late diagnosis of DD as the cause of HC, with a new genetic variant in heterozygosis NP_002285.1:p.Asn242Thrfs*41 in the LAMP2 protein encoding the protein Lysosome-associated membrane glycoprotein 2. This variant of the truncating frameshift mutation has not been previously published, or even identified in controls. No truncating variant of this gene is listed in the database Exome Aggregated Consortium (ExAC). The consequence of this mutation is the creation of a stop codon that causes a premature interruption in the coding of the LAMP2B protein.3,6

Biological diagnosis of DD involves demonstration of normal or high acid maltase activity in combination with muscle biopsies showing large vacuoles filled with glycogen, cytoplasmic degradation products and partial or total absence of LAMP-2 protein in immunohistochemical analysis.1,8 Thus, the diagnosis can be confirmed by molecular analysis of the LAMP2 gene and evaluation of the three isoforms: LAMP 2A, LAMP 2B and LAMP 2C. The LAMP 2B isoform is responsible for metabolic defects, impairing autophagosome-lysosome fusion, leading to heart disease in DD.7

The Wolff-Parkinson-White syndrome is an important diagnosis, especially in women.9 This finding culminated in the indication of the first ablation of the anomalous pathway. In a recent systematic review evaluating 146 patients with DD, while female patients had a predominant pattern of cardiomyopathy alone, as presented in our case report, men more commonly presented the clinical triad of HC, skeletal myopathy and mental illnesses.9

In addition, the reported case had chest pain accompanied by elevated troponin I levels, which raised the suspicion of acute myocarditis. Later, the persistence of elevated troponin levels over the months indicated a false diagnosis of myocarditis. Cardiac resonance has allowed a more accurate evaluation of the site of myocardial hypertrophy, presence of intracavitary thrombi, and recently, presence of fibrosis, which is an important prognostic marker in this group of patients. Myocardial injury and increased troponin may cause myocardial remodeling and explain the progression to a dilated form of myocardial. High levels of troponin I in DD has prognostic value in the clinical decision-making process.10

A recent clinical trial5 studied gene therapy in male patients with DD. This therapy involves a recombinant adeno-associated virus containing the transgene isoform LAMP2B (RP-A501), which will contribute to a better management and understanding of the disease.5

Danon Disease

Danon disease is an uncommon condition, and rarely recognized as a red flag for CH phenocopying. Because there was no specific treatment during the follow-up of the patient reported in this study, several interventions and hospitalizations were necessary over the five years of follow-up. This shows the importance of cardiac surveillance and multidisciplinary approach of this group of patients. Also, telemedicine support allows the maintenance of care despite the COVID-19 pandemic. Finally, genetic testing should be incorporated into clinical practice for congenital heart disease.

Potential Conflict of Interest

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

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There were no external funding sources for this study.

Study Association

This study is not associated with any thesis or dissertation work.

Ethics approval and consent to participate

This article does not contain any studies with human participants or animals performed by any of the authors.

Author contributions

Conception and design of the research: Matos RC, Soares AC, Silva RTB, Mesquita ET. Acquisition of data: Matos RC, Soares AC, Silva RTB, Mesquita ET. Analysis and interpretation of the data: Matos RC, Soares AC, Silva RTB, Mesquita ET. Writing of the manuscript: Matos RC, Soares AC, Mesquita ET. Critical revision of the manuscript for intellectual content: Mesquita ET.
Erratum

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In Case Report “Danon Disease in an Asymptomatic Woman: A Five-Year Follow Up”, with DOI number: https://doi.org/10.36660/ijcs.20210038, published in ahead of print in the journal International Journal of Cardiovascular Sciences, 2022; [online].ahead print, pp.0-0, correct the author’s name “Ricardo Cardoso Cardoso de Matos” to “Ricardo Cardoso de Matos”.

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