Cardiac involvement in Becker-Kiener muscular dystrophy illustrated by cardiac MRI

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Introduction

Becker-Kiener muscular dystrophy (BMD) is as Duchenne dystrophy a X-linked inherited disorder of cardiac and skeletal muscle cells [1]. Cardiac involvement is frequent, mostly unrecognized and yet associated with a high morbidity and mortality [2]. Cardiac MRI (CMR) allows a non-invasive comprehensive analysis of cardiac morphology, function and tissue integrity, the later by late gadolinium enhancement (LGE) [3]. LGE imaging might detect already subclinical cardiac alterations prior to clinical symptoms or echocardiographic findings [4].

Case report

We report about a 39-year male patient with known BMD, who suffered from dyspnea (NYHA II-III), chest pain and arrhythmias. Echocardiography demonstrated a reduced left ventricular (LV) function and a moderate dilatation of the LV. CMR was performed on a 3-Tesla MRI-scanner (Skyra, Siemens, Germany). Axial T2 haste black blood - sequences of the whole thorax, short and long axis CINE and LGE SSFP - sequences were acquired. Additionally, T1 mapping (Molli) was performed in representative short and long axis views.

Global LV function was reduced to 37% (Figure 1), the LV was dilated (LVEDVI 106 ml/m²). LGE showed a predominantly intramyocardial patchy pattern affecting most basal and midventricular septal and inferolateral wall segments (Figure 2). Region-of-interest (ROI) analysis of post-contrast long axis T1-mapping images detected inhomogenously reduced T1 SI values in septal and lateral myocardium (Figure 3), most pronounced in LGE positive areas of the septum.

Discussion

Cardiac involvement is life-limiting in BMD. Varghese and Pennell were the first to describe extensive mid-myocardial late gadolinium enhancement as a hallmark of cardiac disease in BMD [3]. There have also been reports of subendocardial LGE patterns, alluding to the occurrence of myocardial infarctions in these patients [5]. Demonstration of LGE might be a surrogate marker for an impaired clinical outcome [6]. In a series of 15 patients with BMD, Yilmaz and Sechtem showed that CMR with LGE is more sensitive than echocardiography for detection of cardiac alterations [4]. An intramyocardial pattern of LGE positive areas was also seen in the patient presented here. In accordance to the previous reports, mostly septal and inferolateral segments of the LV were affected. Interestingly, also a subepicardial pattern was observed in lateral midventricular parts of the LV, resembling LGE pattern previously reported for myocarditis. A recent large meta-analysis of LGE patterns in non-ischemic cardiomyopathies (NICM), encompassing 9 studies with a total of 1488 patients, proved the negative predictive value for LGE concerning heart failure hospitalization and mortality [7]. Prognostic information of LGE imaging is even increased, when LGE data are quantified [8]. However, the mostly applied methods for

Figure 1. The depiction of left ventricular dilation was possible using long- (1A, 3-chamber-view) and short axis (1B) CINE-sequences.

Figure 2. Intramyocardial late gadolinium enhancement (LGE) in septal and inferolateral LV segments. The long-axis view (2A) demonstrates the predominantly patchy pattern of LGE areas, the short-axis view (2B) a more confluent pattern, including also the subepicardium.
planimetry of LGE areas are time demanding and restricted by intra- and interindividual variabilities.

The new method of calculation of tissue T1 values in pre- and/or post-contrast images might be a more robust and simpler way to establish quantitative data on tissue integrity including the detection of diffuse myocardial fibrosis. Data on t1 mapping in myotonic muscular dystrophy (DM) type 1 and 2 were the first applications of this technique in muscular dystrophies [9]. We saw a similar reduction of post-contrast T1 time in the BMD patient presented in this case report. This likely reflects the presence of diffuse myocardial fibrosis, as we saw also a reduction of t1 time in myocardial segments without LGE.

Florian and Yilmaz used a highly sophisticated approach to calculate t1 mapping results in BMD, generating R1 maps by taking the reciprocal of each T1 map on a pixel-by-pixel basis and ΔR1 maps were computed by subtracting the pre-contrast R1 map from the post-contrast R1 map [10]. Compared to that, we used a simple visual and semiquantitative ROI approach, leading to the same results. In our opinion, ROI analysis is a robust and easy technique to assess changes in myocardial texture in muscular dystrophy.

**Conclusion**

This case report illustrates the feasibility of a comprehensive analysis of cardiac involvement in BMD by cardiac MRI (CMR). The addition of t1 mapping opens the possibility to assess diffuse myocardial damage/fibrosis using the robust and simple approach of myocardial ROI-analysis. The reduction of T1 time is typical in BMD, affecting segments with and without overt fibrosis. In the future, CMR could be applied even in BMD carriers to detect subclinical cardiac involvement like in Duchenne dystrophy [11].

**References**