Evaluation of the differences of myocardial fibers between acute and chronic myocardial infarction: Application of diffusion tensor magnetic resonance imaging in a rhesus monkey model

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Evaluation of the Differences of Myocardial Fibers between Acute and Chronic Myocardial Infarction: Application of Diffusion Tensor Magnetic Resonance Imaging in a Rhesus Monkey Model

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Objective: To understand microstructural changes after myocardial infarction (MI), we evaluated myocardial fibers of rhesus monkeys during acute or chronic MI, and identified the differences of myocardial fibers between acute and chronic MI.

Materials and Methods: Six fixed hearts of rhesus monkeys with left anterior descending coronary artery ligation for 1 hour or 84 days were scanned by diffusion tensor magnetic resonance imaging (MRI) to measure apparent diffusion coefficient (ADC), fractional anisotropy (FA) and helix angle (HA).

Results: Comparing with acute MI monkeys (FA: 0.59 ± 0.02; ADC: 5.0 ± 0.6 x 10^{-4} mm^2/s; HA: 94.5 ± 4.4°), chronic MI monkeys showed remarkably decreased FA value (0.26 ± 0.03), increased ADC value (7.8 ± 0.8 x 10^{-4} mm^2/s), decreased HA transmural range (49.5 ± 4.6°) and serious defects on endocardium in infarcted regions. The HA in infarcted regions shifted to more components of negative left-handed helix in chronic MI monkeys (-38.3 ± 5.0°–11.2 ± 4.3°) than in acute MI monkeys (-41.4 ± 5.1°–53.1 ± 3.7°), but the HA in remote regions shifted to more components of positive right-handed helix in chronic MI monkeys (-43.8 ± 2.7°–66.5 ± 4.9°) than in acute MI monkeys (-59.5 ± 3.4°–64.9 ± 4.3°).

Conclusion: Diffusion tensor MRI method helps to quantify differences of mechanical microstructure and water diffusion of myocardial fibers between acute and chronic MI monkey’s models.

Keywords: Rhesus monkey; Myocardial fiber; Myocardial infarction; Helix angle; Diffusion tensor MR imaging; DTI

INTRODUCTION

Previous studies have confirmed that the deterioration of cardiac mechanical function originates from micro-alterations of myocardial fibers after myocardial infarction (MI) (1, 2). Compared with the conventional histological method, diffusion tenor magnetic resonance imaging (DT-MRI) is a highly sensitive approach to evaluate myocardial fibers in a 3D manner (3-5). However, in-vivo DT-MRI for myocardial fibers is limited by the image distortion that is induced by the cardiac motion and the amount of time required to complete the operation (6, 7). Alternatively, ex-vivo DT-MRI for animal models provides some valuable
hypothesized that the diffusion and mechanical differences in myocardial fibers between acute and chronic MI could be quantified and differentiated using the ex vivo DT-MRI method.

The objective of this study was to quantify the characteristics of the myocardial fibers during acute and chronic MI, and subsequently identify the differences in myocardial fibers between acute and chronic MI in rhesus monkeys using the DT-MRI method, in order to understand the corresponding changes in patients with acute MI (16).

MATERIALS AND METHODS

Experiments

All experiments were performed in accordance with the
guidelines set by the U.S. National Institute of Health and the regulations for the humane care of laboratory animals of our university. The study protocol was approved by the Institutional Review Committee on Animal Care of the provincial government.

Six adult rhesus monkeys (3 males and 3 females, aged 3 years) were randomized into the acute MI (n = 3) or the chronic MI (n = 3) group. All monkeys were anesthetized deeply by pentobarbital and underwent permanent suture ligation of the left anterior descending coronary artery. Monkey’s chests were closed and fed for 1 hour or 84 days, named as MI-1 hour (n = 3) or MI-84 day (n = 3), respectively. Delayed enhancement magnetic resonance imaging (DE-MRI) was used to confirm infarcted regions in the left ventricular in vivo (Fig. 1). After two hours of completing DE-MRI, all infarcted monkeys were sacrificed and their hearts were excised under deep surgical anesthesia. All excised hearts were perfusion-fixed by a 4% paraformaldehyde solution and stored in a 1% paraformaldehyde solution.

Before DT-MRI scanning, Fomblin, special lubricating oil (per-fluororated poly ethers), was injected into four chambers of each excised heart to reduce susceptibility artifacts in myocardial margins during DT-MRI scanning, and then, each excised heart was sealed in a plastic container filled with Fomblin.

With the heart container in a 7T MR (Bruker BioSpec 70/30, Chengdu, China) scanner, DT-MRI was performed to evaluate the myocardial fibers after MI. The echo-planar imaging-based diffusion tensor sequence was applied to acquire a single non-weighted (B0) image and diffusion images in 30 gradient directions. The scanned parameters were listed as follows: repetition time/echo time = 12000/32 ms, matrix = 100 x 100, field of view = 50 x 50 mm, slice thickness = 0.8 mm, flip angle = 90°, segment = 16, number of excitation = 25 and b value = 1300 ms/μm². After DT-MRI, the hearts were sectioned and fixed, and infarcted regions and orientations of myocardial fibers were observed by Masson staining.

Data Analysis

Before calculating the parameters of DT-MRI, the diffusion images in 30 gradient directions were re-aligned with the B0 image to correct the movement errors due to mild scanner vibration (17-19). This step was taken with in-house programs that were encoded using the MATLAB platform (MathWorks, Natick, MA, USA). Afterwards, the singular value decomposition method was employed to calculate three eigenvectors and the corresponding eigenvalues of each voxel in the myocardial tissues, which were essential for the subsequent analysis of the various characteristics of the myocardial fibers.

The FA values and ADC values were calculated by voxel-based three eigenvalues to quantify the diffusion characteristics of the myocardial fibers. This step was performed using the diffusion toolkit package (Massachusetts General Hospital, Boston, MA, USA). The voxel-based tensors and the helix angle (HA) were

| Group                | Number | FA       | ADC       | Range of HA                       |
|----------------------|--------|----------|-----------|-----------------------------------|
| MI-1 hour (infarcted) |        |          |           |                                   |
| Mean                 | 0.59 ± 0.02 | 5.0 ± 0.6 x 10^-4 mm²/s | 94.5 ± 4.4° (41.4 ± 5.1°–53.1 ± 3.7°) |
| 1                    | 0.61   | 4.6 x 10^-4 mm²/s | 98.6° (-42.2°–56.4°) |
| 2                    | 0.59   | 5.3 x 10^-4 mm²/s | 95.2° (-46.0°–49.2°) |
| 3                    | 0.58   | 5.8 x 10^-4 mm²/s | 89.8° (-36.0°–53.8°) |

| MI-1 hour (remote)   |        |          |           |                                   |
| Mean                 | 0.62 ± 0.03 | 4.8 ± 0.8 x 10^-4 mm²/s | 124.4 ± 3.2° (-59.5 ± 3.4°–64.9 ± 4.3°) |
| 1                    | 0.65   | 3.8 x 10^-4 mm²/s | 127.7° (-61.6°–66.1°) |
| 2                    | 0.59   | 5.3 x 10^-4 mm²/s | 121.4° (-61.3°–60.1°) |
| 3                    | 0.60   | 5.2 x 10^-4 mm²/s | 124.0° (-55.5°–68.5°) |

| MI-84 days (infarcted) |        |          |           |                                   |
| Mean                 | 0.26 ± 0.03 | 7.8 ± 0.8 x 10^-4 mm²/s | 49.5 ± 4.6° (-38.3 ± 5.0°–11.2 ± 4.3°) |
| 1                    | 0.25   | 8.2 x 10^-4 mm²/s | 49.0° (-42.5°–6.5°) |
| 2                    | 0.3    | 6.9 x 10^-4 mm²/s | 54.6° (-39.6°–14.8°) |
| 3                    | 0.24   | 8.4 x 10^-4 mm²/s | 45.2° (-32.8°–12.4°) |

| MI-84 days (remote)  |        |          |           |                                   |
| Mean                 | 0.59 ± 0.02 | 4.9 ± 0.7 x 10^-4 mm²/s | 110.3 ± 3.6° (-43.8 ± 2.7°–66.5 ± 4.9°) |
| 1                    | 0.58   | 5.4 x 10^-4 mm²/s | 106.3° (-43.2°–63.1°) |
| 2                    | 0.62   | 4.1 x 10^-4 mm²/s | 113.5° (-41.4°–72.1°) |
| 3                    | 0.58   | 5.3 x 10^-4 mm²/s | 111.1° (-46.7°–64.4°) |

ADC = apparent diffusion coefficients, FA = fractional anisotropy, HA = helix angle, MI = myocardial infarction
calculated by voxel-based primary eigenvectors to quantify the mechanical and microstructural characteristics of the myocardial fibers. This step was taken with in-house programs that were encoded using the MATLAB platform (MathWorks, Natick, MA, USA).

RESULTS

The FA, HA, and ADC values of each monkey and the mean values were listed in Table 1. Figure 2 illustrated FA and ADC maps of the infarcted slices in the MI-1 hour and MI-84 day monkeys. In the group of MI-1 hour monkeys, compared with the remote regions (FA: 0.62 ± 0.03, ADC: 4.8 ± 0.8 \( \times 10^{-4} \) mm\(^2\)/s), the MI regions (red arrow) showed slightly decreased FA (0.59 ± 0.02) and increased ADC values (5.0 ± 0.6 \( \times 10^{-4} \) mm\(^2\)/s), which are involved in early intracellular edema of the myocytes. In contrast, in the group of MI-84 days monkeys, compared with the remote regions (FA: 0.59 ± 0.02, ADC: 4.9 ± 0.7 \( \times 10^{-4} \) mm\(^2\)/s), the MI regions (red arrow) showed noticeably decreased FA (0.26 ± 0.03) and increased ADC values (7.8 ± 0.8 \( \times 10^{-4} \) mm\(^2\)/s), which indicated severity of necrotic myocytes.

Figure 3 is the HA and tensor map of the infarcted slices in the MI-1 hour and MI-84 days monkeys. In the group of MI-1 hour monkeys, compared with the remote regions (HA: 124.4 ± 3.2°), there were mild changes in the HA transmural ranges (94.5 ± 4.4°) of the infarcted region (black circle) and the primary orientations of the tensor. The tensors became slightly thinner in the endocardium but not in the epicardium. In contrast, the HA values (remote regions: 110.3 ± 3.6°, infarcted regions: 49.5 ± 4.6°) in the MI-84 days monkeys were not homogeneous. Extensive tissue defects were observed in the endocardium. The decreased HA indicated the reduced mechanical function.

Fig. 2. Diffusion maps of myocardial fibers. Regions pointed by red arrows are MI regions. ADC = apparent diffusion coefficients, FA = fractional anisotropy, MI = myocardial infarction.
and the increased electrophysiological risk in the infarcted regions.

Figure 4 enables the comparison of the tensor map with the Masson staining image. The MI-1 hour panel showed that the density of the myocardial fibers decreased slightly. The tensor map also showed a slightly decreased arrangement and density of the tensors in the myocardial fibers. The Masson staining image in the MI-84 days panel showed an expansive area of blue collagens and myocardial fibers in the blue collagens that are sparsely distributed. The tensor map showed some defects of the tensors in the endocardium.

Figure 5 illustrated the transmural range of HA from the epicardium to the endocardium on the anterior and posterior walls in the MI-1 hour and MI-84 days monkeys. Compared with the MI-1 hour monkeys (infarcted: -41.4 ± 5.1°–53.1 ± 3.7°, remote: -59.5 ± 3.4°–64.9 ± 4.3°), the infarcted regions in the MI-84 days monkeys showed a significantly decreased HA transmural range and a significant shift of the HA transmural range from the double-helix to the left-handed helix (-38.3 ± 5.0°–11.2 ± 4.3°), but the remote regions showed a significant shift of the HA transmural range from the double-helix to the right-handed helix (-43.8 ± 2.7°–66.5 ± 4.9°). The changed HA transmural range indicated decreased wall torsion in the infarcted regions and mechanical compensation in the remote region.

**DISCUSSION**

In this study, the DT-MRI method was applied successfully to quantify the characteristics of myocardial fibers in rhesus monkeys with acute and chronic MI. Our results confirmed that there were differences in myocardial tensor and diffusion between the acute and chronic MI monkey models. Moreover, our findings provided a more comprehensive understanding of the early changes of myocardial fibers in patients with acute MI.

**Diffusion Parameters: FA and ADC Values**

The FA value is considered as the longitudinal-to-transverse aspect ratio of the myocytes (6, 20). During the MI-1 hour period, many spherical cells such as inflammatory cells, swollen myocytes and myofibroblasts began to accumulate in the infarcted regions, which slightly reduced the FA values in the infarcted regions due to the slight reduction of the longitudinal-to-transverse ratio of the myocytes. When the time was extended to MI-84 days, the myocardial fibers in the scar tissues became disarrayed, and extensive collagens replaced the extracellular matrix in the infarcted regions, which resulted in the severe reduction of the FA values in the infarcted regions. Our findings were consistent with those from a previous histological study; more severe MI revealed lower FA values in the infarcted regions (12, 21).

The ADC value was used to quantify the balance of free water between the intracellular and extracellular space volumes (6). During the MI-1 hour, the increased lactic acid damaged the protein in the infarcted region, and the combined water in the protein became free water; subsequently, the elevated osmotic pressures caused by the intracellular Na+ accumulation caused the free water...
to enter into the intracellular space due to dysfunctions of the Na+/K+ channels (22, 23). Compared with the remote regions, the infarcted region showed a slight increment of free water in the intracellular space, which mildly increased the ADC values in the infarcted regions. In the MI-84 days, the extracellular space in the infarcted regions was increased significantly since the aggregations of fibroblasts, neutrophils, and macrophages did not fill the extracellular space created by the necrotic myocytes after MI (6). Accordingly, the significantly increased ADC values were attributed to the increased extracellular space in the infarcted regions.

Fig. 4. Comparison between tensor maps and Masson’s staining images. On MI-1 hour panel, orientations of myocardial fibers remain consistent. Tensor map also reveals similar arrangements and density of tensors in myocardial fibers. On MI-84 days panel, Masson’s staining image shows large area of blue collagens, myocardial fibers in blue collagens are sparsely distributed, and orientation of myocardial fibers are inhomogeneous. Tensor map displays some tissue defects of tensors in endocardium and tensors are irregularly arranged. MI = myocardial infarction

Fig. 5. Remodeling process of HA transmural range from epicardium to endocardium on anterior and posterior walls during MI-1 hour and MI-84 days monkeys. HA transmural range in infarcted region (anterior) is decreased but shift from double-helix toward left-handed helix during chronic MI. However, HA transmural range in remote (posterior) region shows increased shift from double-helix toward right-handed helix during chronic MI. HA = helix angle, MI = myocardial infarction
Mechanical Parameters: Tensors and HA Values

The voxel-based tensors are the important mechanical microstructures in the myocardial fibers because the wall torsion depends on the movements of the multi-orientation tensors (16, 24). In the MI-1 hour, the early stage of myocardial ischemia resulted in moderate myocardial edema, but the capillary wall retained its structural integrity, and the myocytes did not undergo extensive apoptosis. In the MI-84 days, injury on the infarcted wall was transmural from the endocardium to the epicardium. Some tissue defects of the myocardial tensors occurred in the endocardium of the infarcted wall. The scar and collagens in the residual wall resulted in the irregular arrangements of the tensors, which significantly increased the electrophysiological risk (14).

The transmural range of HA was widely applied to quantify the orientations of myocardial fibers from the epicardium to the endocardium (25-31). In the MI-1 hour, the change in the transmural range of HA was moderate as the orientations of the tensors remained essentially unchanged in the infarcted region. In the MI-84 days, the transmural range of HA was significantly reduced due to tissue defects of the tensors on the endocardium. The tissue defects on the endocardium resulted in the reduction of the positive right-handed helix in the infarcted regions, which revealed the shift from the transmural double-helix to the residual negative left-handed helix on the epicardium. However, the remote regions revealed the shift from the transmural double-helix to the reduced negative left-handed helix on the epicardium. Our findings in rhesus monkeys were consistent with the findings of previous studies that were conducted using other animals (6, 12, 32-34). We proposed that the HA shifts during MI-84 day were indicative of decreased wall torsion and contractile efficiency in the infarcted and mechanical compensation in the remote region (35).

Moreover, the FA values, ADC values or transmural range of HA in the infarcted regions were approximately 0.26, 0.00078 mm$^2$/s or $-38 \pm 11^\circ$ in the MI-84 days monkeys, respectively, and the values were close to those in the MI patients (0.26, 0.000824 mm$^2$/s or $-36 \pm 10^\circ$) (6, 12, 21, 32-34). Thus, the mild changes of water diffusion and myocardial tensors in the MI-1 hour monkeys could contribute to the understanding and evaluation of these parameters in patients during the early stage of acute MI.

Limitations

More monkeys should be recruited in our future study. Formalin fixation decreased ex-vivo diffusion of myocardium (36-38). The effect of temperature on ex-vivo diffusion of myocardium might be increased due to the increased coil temperature caused by the long scanning period.

In conclusion, our study confirmed that the DT-MRI method could quantify and identify the acute and chronic MI monkey models by comparing the mechanical microstructure and water diffusion of myocardial fibers. According to the similar changes between monkeys and chronic MI patients, the characteristics of myocardial fibers in the acute MI monkeys could contribute to the understanding and evaluation of these parameters in patients during the early stage of acute MI.

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