Target Audience: Scientists/clinicians interested in MRI of the myocardium.

Purpose: The area of myocardium adjacent to an infarct (border zone) plays a major role in the development of mechanical and electrical dysfunction as the heart remodels. Intermediate signal on late gadolinium enhancement (LGE) has been used to delineate the infarct periphery and border zone. However, the microstructural properties of the border zone in the human heart in vivo have not yet been characterized. Here we used in vivo diffusion tensor MRI (DTI) and LGE in patients with recent myocardial infarctions to elucidate the microstructural properties of the infarct border zone.

Methods: Patients with acute myocardial infarction (MI, n=3) were scanned prior to discharge on a clinical 1.5T scanner with a diffusion-encoded stimulated echo EPI sequence and the following parameters: resolution 1.75x1.75x8 mm$^3$, b-value of 500 s/mm$^2$, 6 diffusion encoding directions and 8 averages. Imaging was performed in the diastolic sweet spot of the cardiac cycle to mitigate the effect of strain. Late gadolinium enhancement (LGE) was performed after the injection of 0.2 mmol/kg of Gd-DTPA using an inversion recovery gradient echo sequence. Infarcted myocardium by LGE was defined by a signal intensity > 2 standard deviations (SD) above normal myocardium. The infarct was further subdivided into the infarct core (>3 SD above normal) and the infarct periphery (2-3 SD above normal). The border zone was defined by signal between 1-2 SD above normal. The diffusion tensor field was determined and diagonalized to yield the principal (e$_1$/\lambda_1$), secondary (e$_2$/\lambda_2$), and tertiary (e$_3$/\lambda_3$) eigenvectors/values. Mean diffusivity (MD), fractional anisotropy (FA), helix angle (HA), and propagation angle (PA) were calculated as previously described. In addition, HA variance at several transmural depths was measured. Fiber tracts were constructed by integrating the primary eigenvector field into streamlines using a 5th order adaptive Runge-Kutta approach and color coded by HA. Values in the infarct, border, and remote zones were compared using ANOVA.

Results: A total of 16 short axis LGE images were segmented and analyzed. A representative LGE image before and after segmentation is shown in Figure 1A-B. MD, FA, HA, and PA maps at the corresponding slice location are shown in Figures 1C-F. MD, PA, and HA variance in the border zone were significantly higher than that of normal myocardium but also lower than in the infarct zone. Likewise, FA in the border zone was lower than that of normal myocardium but higher than in the infarct. Average values are plotted in Figure 2A-D. Fiber architecture in the remote zone showed the characteristic crossing helical pattern of subendocardial (blue) and subepicardial (yellow-red) fibers (Fig. 2E). Fiber architecture in the border zone was markedly perturbed and heterogeneous (Fig. 2F). While coherent tracts could be resolved in some portions of the border zone, large gaps with no coherent fiber tracts were seen. In addition, the orientation of the tracts in the subendocardium and subepicardium was similar and the characteristic crossing helical pattern was lost. Severe microstructural disarray was seen in the infarct zone with the near complete loss of normal fiber orientation (Fig. 2G).

Conclusion: The border zone has a highly complex and heterogeneous microstructure that can be resolved with in vivo DTI. This could play an important role in clinical care, where knowledge of the specific microstructure of the border zone could be of major value.