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Poster presentation

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## Evolution of edema, hemorrhage and microvascular obstruction after acute myocardial infarction

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#### Introduction

In acute myocardial infarction (AMI), the no-reflow phenomenon is caused by ischemia-induced microvascular injury/obstruction and has been correlated with adverse remodeling. The severity of the initial ischemic insult may also lead to intramyocardial hemorrhage. Alongside, intracellular and interstitial edema is a consistent feature of AMI and has been associated with the salvageable areat-risk. The (in-vivo) evolution of these processes throughout infarct healing is not well-characterized but is important in grading severity and evaluating treatment strategies, potentially improving clinical outcome.

#### **Purpose**

To characterize the time course of edema (T2), hemorrhage (T2\*) and microvascular obstruction (MVO) in porcine myocardium following AMI and observe the relative resolution of these pathophysiological mechanisms.

### **Methods**

7 pigs underwent MRI before LAD infarction (control) with subgroups studied at 2,7,14, and 30-42 days post-infarction. Histology was performed upon sacrifice at either Day 14 (n = 3) or Day 30-42 (n = 4). Imaging was performed on a 3 T MRI scanner (MR 750, GE Health-care). A previously validated T2-prepared spiral sequence was utilized for T2 quantification and T2\* was determined using a multi-echo gradient-echo acquisition. An early (~3 min) contrast-enhanced (CE) IR-GRE sequence

was used for infarct/MVO delineation. Diastolic-wall-thickness (DWT) was measured from CINE-SSFP imaging.

## **Results**

Figure 1 demonstrates T2, T2\* maps and early CE images for an anterio-septal infarct in a short-axis slice for a representative animal at three time points. T2-maps represent edematous changes (bright regions), T2\*-maps indicate hemorrhage (dark regions) while CE images delineate MVO (signal voids within infarct). Figure 2 shows the cumulative time course of T2, T2\* and DWT within the infarct. T2 was indistinguishable from control at day 2 (p = 0.38) while the T2 elevation beyond week 1 was statistically significant (p < 0.05). T2\* was reduced up to week 1 as a result of hemorrhage and its normalization at week 4 coincided with resolution of MVO. DWT was significantly increased at day 2 (7.5 vs 5.3 mm, p = 0.06) suggesting increased tissue water content while it fell below control values at week 6 (4.3 mm, p = 0.003) indicating scar formation.

#### Conclusion

Post-infarct remodeling is a complex process and comparison with remote myocardium is equally important. In this respect quantitative T2 and T2\* mapping techniques are potentially more specific than intensity measures in single images. Edema and hemorrhage have counter-acting effects on T2, hence care should be taken while evaluating day 2. Our study demonstrates that multi-factorial

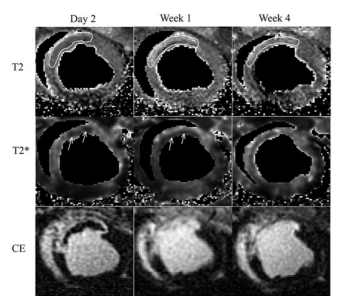


Figure I
At day 2 in this animal, T2 elevation usually associated with edema was not apparent in the infarct zone (39.2 ms vs 39.1 ms contrl); however DWT was increased by 34% suggesting edematous swelling.

Lower T2\* (arrows) indicated presence of hemorrhage (18.5 ms vs 34.2 ms while the CE image showed a large MVO. At Week I, T2 was elevated in most of the infarct (51.1 ms) with reduced T2\* (20.5 ms) indicating diffuse hemorrhagic by-products. CE image showed only a slight MVO. By week 4, hemorrhage/MVO were resolved.

MR-based parameters, acquired in a longitudinal fashion, can be employed to assess the evolution of myocardial infarction.

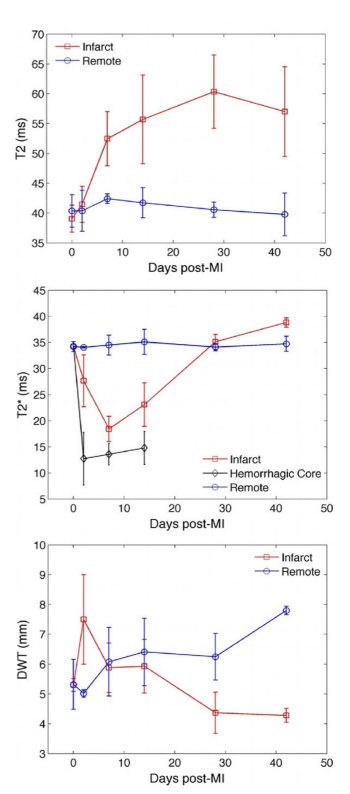


Figure 2
Plots demostrate longitudinal fluctuations in T2, T2\*
and DWT in infarct zone compared to remote myocardium averaged over all animals