INTRODUCTION: Late Gadolinium Enhancement (LGE) CMR is the gold standard of imaging irreversible damage in myocardial infarction. However, limitations include invasive contrast administration and sensitivity to accurate choice of Inversion Time (TI). Alternative non-invasive quantitative mapping of the myocardium has been demonstrated to distinguish pathophysiological changes in acute myocardial infarction at 1.5T [1]. We present preliminary experiences of experimental $T_1$ [2, 3] and $T_2$ [4] mapping techniques at 3T to distinguish areas of affected from unaffected myocardium.

METHODS: Clinical Material and Methods: 4 patients with a first acute myocardial infarction (age 53±10 years; 3 males) underwent CMR imaging at 3T (TRIO, SIEMENS). LGE images were obtained 24-48 hours post acute infarct [5]; $T_1$ and $T_2$ maps were obtained 5-17 days after the ischemic event. Pilot $T_1$-maps using the novel ShMOLLI sequence (a shortened version of MOLLI [1]) and $T_2$-maps [4] at a single representative slice were generated. ShMOLLI was implemented as 3 IR experiments split over 9 heartbeats (separated by only one heartbeat) to collect 5+1+1 SSFP images with varying TI (typically 110-5000ms, TE=1.1ms, FOV=360x280mm, matrix 192x144, interpolation=2, pixel size≈0.9mm). ShMOLLI samples from the second and third IR are taken into account only if the estimated $T_1$ is shorter than the R-R interval, and they improve nonlinear fit. The nonlinear fitting was implemented in C++ directly in the scanner reconstruction pipeline utilizing parallel processing with images available for viewing directly on console immediately after acquisition. $T_2$ maps were generated from a series of 5 $T_2$ prepared images (TE=0, 32, 55, 78, 100ms), also reconstructed directly on the scanner. Imaging parameters: $T_2$-prepared with single shot SSFP acquisition, TR/TE=313/1.04ms, flip angle 48º, FOV 370x270, acquisition matrix 128x116 interpolation=2, pixel size≈1.1mm. Post-processing involved manual segmentation of the myocardium followed by calculation of the distribution of $T_1$ and $T_2$ relaxation times. These were fitted into 2 component Gaussians (See Fig. 1) to separately estimate the distributions for affected and unaffected myocardium.

Phantom verification: Both ShMOLLI and $T_2$ prep-SSFP were validated using separate sets of 50ml Agarose+NiCl gel phantoms [6] with $T_2$=60ms and $T_1$=70-2300ms (for ShMOLLI) and $T_1$=900-1500ms, $T_2$=30-100ms (for $T_2$Prep-SSFP). For $T_1$ reference we used a spin echo sequence with TI=33, 100, 300, 900, 2700, 5000 ms, TE/TR=6.3ms/10s. For $T_2$ estimation we used Spin Echo with TR=100ms and TE=6.3, 12, 24, 40, 80, 150, 250, 500ms. Reference images were fitted offline using non-linear methods separately for $T_1$ and $T_2$. The average estimates of the phantoms were used as baseline to obtain empirical correction for the in-vivo measurements.

RESULTS&DISCUSSION: Phantom study showed that $T_1$-mapping using ShMOLLI underestimates $T_1$ values by ≈4%. $T_2$-mapping using $T_2$-prep has relatively poor metrological properties as identified by the empirical relationship: $T_2$prep=0.8*$T_{2\text{reference}}+20\text{ms}$; (R$^2$=0.9). These corrections have been used to present the measured relaxation times in Table 1. The application of the methods in clinical cases was easy due to short imaging times. $T_2$ maps were of good quality and showed distinctly separate distribution peaks within the myocardium (Fig.1A). $T_2$ maps were less robust but, with the exception of case #3 which was affected by a large artefact in the anterolateral wall, it was possible to assess the entire myocardial rim and obtain similar bimodal histograms. LGE lesions overlapped with areas characterized by increased relaxation times. These were fitted into 2 component Gaussians (See Fig. 1) to separately estimate the distributions for affected and unaffected myocardium. The width of estimated peaks was about 4-5% of the average estimates. This compares favourably to the estimated relative changes in the relaxation times (15-25% for $T_1$; 26-60% for $T_2$), making relaxation maps a good target for automated objective lesion segmentation.

Conclusion: $T_1$ and $T_2$ relaxation times in infarcted myocardium demonstrate distinctively separate distribution peaks that co-localise with LG enhanced regions of damage. While the underlying pathophysiological phenomena mirrored in relaxation properties remain to be established, this observation potentially paves the way to objective lesion segmentation without the need for contrast agents.

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