

WORKSHOP PRESENTATION

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Free-breathing myocardial T_1 mapping using magnetization-prepared slice interleaved spoiled gradient echo imaging

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Background

Quantitative myocardial T_1 mapping and extracellular volume fraction (ECV) show promise for non-invasive assessment of cardiomyopathies. Most available T_1 mapping sequences use a single slice breath-hold acquisition with balanced steady state free precession

(b-SSFP) readout [1]. However, b-SSFP readout is sensitive to B_0 field inhomogeneity and is potentially T_2 dependent [1]. In this study, we sought to investigate the feasibility of a free breathing multi-slice T_1 mapping sequence using slice-interleaved spoiled gradient echo (GRE) imaging.

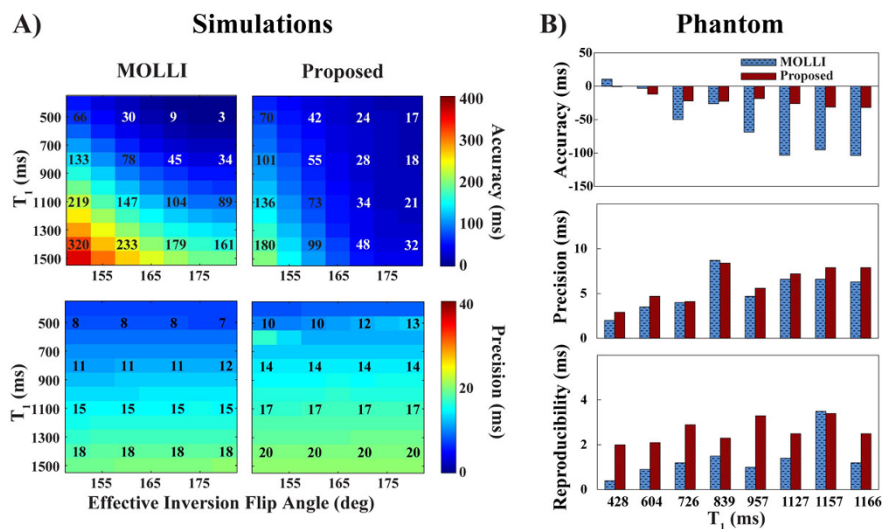


Figure 1 Accuracy, precision and reproducibility of the proposed sequence obtained in Monte Carlo simulation (20,000 repetitions, fixed T_2 of 50 ms, SNR corresponding to 50 in the ∞ image) (a) and phantom experiments (set of vials with NiCl₂ doped agarose, 15 repetitions of the sequence) (b). Results were compared to the MOLLI (5-(3)-3 scheme) sequence. Accuracy was measured in each vial as the difference between spin echo T_1 measurements and the average T_1 over all 15 repetitions. Precision was measured in each vial as the average (over all 15 repetitions) of the standard deviation of T_1 within a vial. Reproducibility was measured in each vial as the standard deviation (over all 15 repetitions) of the mean T_1 within a vial. Improved accuracy and similar precision were achieved using the proposed sequence in both simulations and phantom experiments. T_1 mapping reproducibility was slightly decreased with the proposed sequence.

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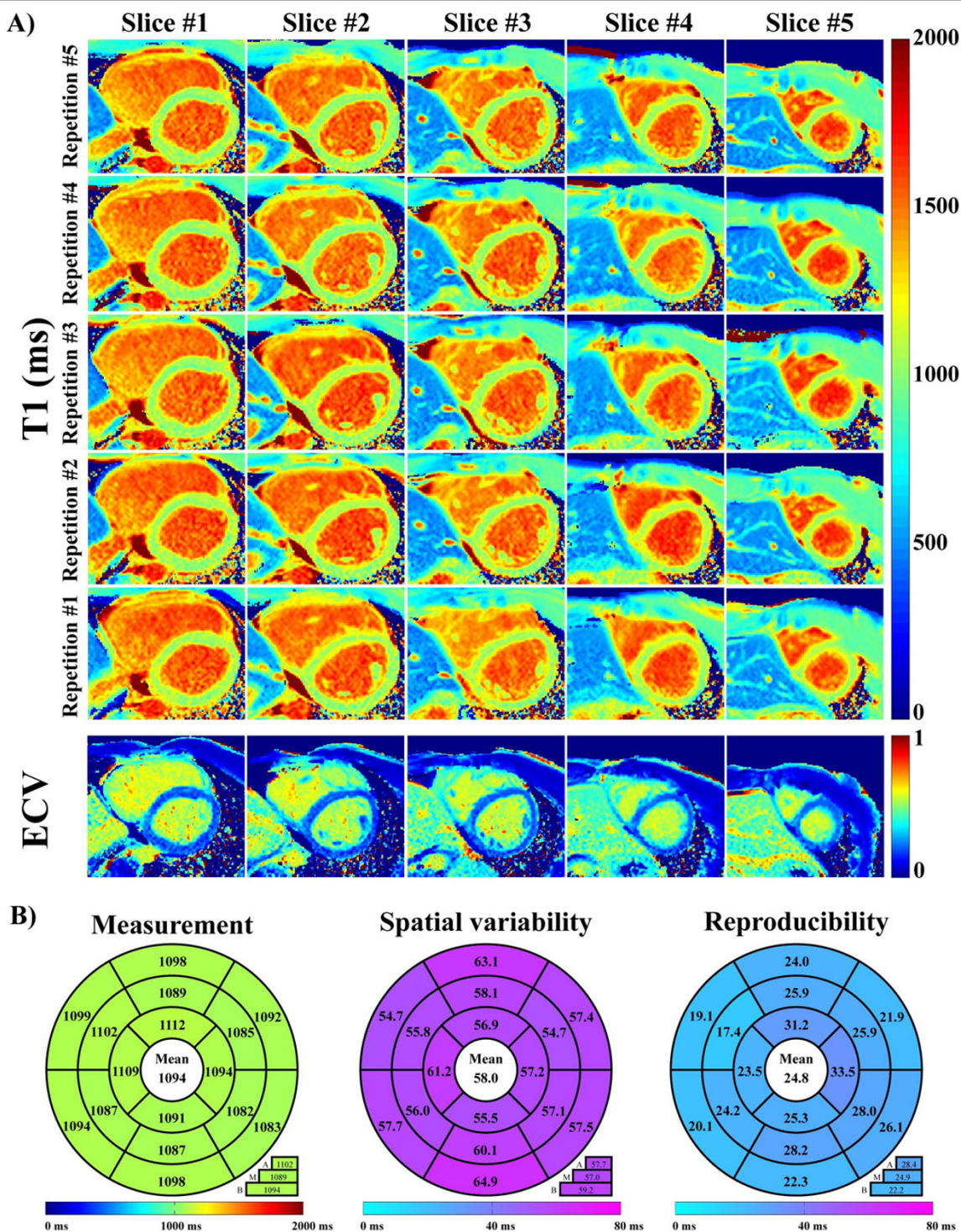


Figure 2 In-vivo native T_1 and ECV mapping using the proposed sequence. Example of multi-slice T_1 maps and ECV maps obtained in one healthy subject is shown in (a). Homogeneous T_1 map quality was achieved in all slices for all five repetitions. Homogeneous ECV map quality was also observed through all slices. Native T_1 measurements, spatial variability, and reproducibility obtained using the proposed sequence, are reported in average over all subjects in (b). Each metric was quantified using a 16 myocardial segment model in all subjects by analysis of the three mid-ventricular slices. Spatial variability was measured for each segment as the average (over the five repetitions of all subjects) of the standard deviation of T_1 measurements within that segment. Reproducibility was measured for each segment as the average (over all subjects) of the standard deviation (over the 5 repetitions) of the mean T_1 time of that segment.

Methods

The proposed sequence used multiple inversion recovery (IR) experiments. In each IR experiment, a non-selective inversion pulse is applied and followed by the acquisition of 5 slices over the next 5 heart beats, and 3 rest cycles [2]. This IR experiment is repeated 5 times using different slice orders to obtain signal samples at TI, TI + 1 RR, TI + 2 RR, TI + 3 RR, TI + 4 RR. This block of 5 IR experiments is finally repeated using a different TI value. The fully recovered longitudinal magnetization is also initially acquired for each slice without any IR pulse (∞ image). Respiratory motion was corrected using prospective slice tracking and retrospective image registration. ECG-triggered single shot acquisitions were used with GRE readout (TR/TE/ α =4.3/2.1ms/10°, FOV=280×272 mm², voxel size=2×2 mm², slice thickness=8 mm, 5 slices, 43 phase-encoding lines, linear ordering, 10 linear ramp-up pulses, SENSE factor=2.5, half Fourier=0.75, bandwidth=382Hz/pixel). For comparison, MOLLI [3] was acquired with a b-SSFP readout and similar parameters (except TR/TE/ α =2.6/1.3ms/70°, 1 slice, bandwidth=1785 Hz/pixel). Imaging was performed on a 1.5 T Philips scanner. T₁ accuracy, precision, and reproducibility were evaluated in simulations and phantom. In-vivo spatial variability and reproducibility of native T₁ mapping was measured in 11 healthy adult subjects (35±21y, 4 m), imaged 5 times with each sequence. Three of these subjects were also imaged at ~15min after contrast injection to demonstrate the feasibility of ECV mapping.

Results

The proposed sequence provided improved accuracy and similar precision than MOLLI in both simulation and phantom experiments (accuracy: p=0.01; precision: p=0.16). MOLLI was more reproducible in phantom (p<0.001). In-vivo, the proposed sequence yielded higher native T₁ times than MOLLI (1094±24ms vs. 1010±27ms, p<0.001) with similar spatial variability (58±7ms vs. 61±9ms, p=0.44) and reproducibility (25±9ms vs. 17±8ms, p=0.15). ECV measurements were 0.21±0.01 using the proposed sequence.

Conclusions

Free breathing multi-slice T₁ mapping using a magnetization-prepared slice interleaved spoiled GRE imaging is feasible and yields similar in-vivo precision/reproducibility as MOLLI but with improved accuracy. In addition, the proposed sequence allows simultaneous imaging of 5 slices within free-breathing in 100 sec.

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References

1. Kelmann: *JCMR* 2014.
2. Weingärtner: *MRM* 2014.
3. Messroghli: *MRM* 2004.

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