Hyperpolarized [1,4-13C2]Fumarate is a probe of necrosis in myocardial infarction

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Synopsis

Previous work has shown that hyperpolarized [1,4-13C2]fumarate is a probe of cellular necrosis. We demonstrate here that the ratio of cardiac hyperpolarized malate to fumarate is increased by a factor of ∼82 one day after cryoinduced myocardial infarction in rats, decreasing to an ∼30-fold increase one week after injury. We additionally image this injury with a novel spiral multiband pulse sequence. Hyperpolarized fumarate therefore forms a sensitive probe of myocardial injury in vivo, and could form a clinical monitor of cellular damage and necrosis after infarction.

Purpose

Myocardial infarction (MI) is characterized by cellular necrosis and the subsequent cascade of structural and functional adaptations to it. The accurate and reproducible quantification of the region of necrosis immediately following myocardial infarction has been identified as a clinical requirement, and techniques such as late gadolinium imaging or T2-weighted imaging aim to infer the region of necrosis indirectly.[1] At present, there is no clinically validated mechanism for detecting necrosis directly in vivo. The injection of hyperpolarized [1,4-13C2]fumarate and its subsequent conversion to [1,4-13C2]malate has probed cellular necrosis in vivo in tumours[2] and following acute kidney injury.[3] We therefore investigated the use of hyperpolarized [1,4-13C2]fumarate in a rodent model of myocardial infarction.

Methods

Hyperpolarization: [1,4-13C2]Fumaric acid (3.23 mmol; Cambridge Isotopes) was dissolved in 8.74 mmol of DMSO containing 11.48 μmol of a trityl radical (AH111501; GE Healthcare) and 0.48 μmol of a gadolinium chelate (Dotarem, Guerbet). A 40 mg aliquot was polarized per experiment (prototype hyperpolarizer, 45 min at 94 GHz) before dissolution (in 6 ml NaOH, final concentration 20 mM). Infusion (2 ml) was via a tail vein cannula over 20 s following dissolution.

Myocardial Infarction: Six female Wistar rats (~200 g, Harlan) were divided into two groups (MI or control, n = 3), and subject to cryoinduced myocardial infarction.[4] Control animals were subject only to pericardial removal. Pre-operative and postoperative analgesia was provided. All experiments were performed with appropriate ethical review.

Spectroscopy: An actively detuned transmit/surface receive coil was used (72 mm 1H/13C proton/carbon birdcage transmit with 40 mm two-channel 13C surface receive; Rapid Biomedical GMBH, Rimpar, Germany) as described previously.[5] A 5 M 13C-urea phantom was included as a per-animal FA and frequency reference. Slice-selective spectra (200 μs gauss, 20 mm thick, TR = 1 s, bandwidth 10 kHz, FA 10°) were acquired from the heart in end-systole. Cardiac function was assessed by CINE.[6] Spectra were summed for 60 s following the appearance of the [1,4-13C2]fumarate peak, and quantified with AMARES, with the reported malate:fumarate ratio being that of total visible malate to fumarate.

Multiband Imaging: A hybrid multiband spatial-spectral RF excitation pulse[7] was designed to simultaneously excite [1,4-13C2]fumarate at a 4° FA, and both [1,4-13C2]malate resonances at ∼20°, and a slab thickness of 20 mm. A multi-echo spiral readout[8] was constructed to maximise the effective number of signal averages over all chemical species in the subsequent reconstruction, with FID acquisition every seventh echo (TE = 2.1, 3.8, 5.5, 7.2, 8.9, 10.6, 12.3 FID 2.1 ms). The spiral trajectory was designed with a nominal FOV of 80 × 80 mm², readout bandwidth of 250 kHz, TR = 1 RR interval, nominal acquisition in-plane resolution 2 × 2 mm². The pulse sequence is shown in Fig. 1. The k-space trajectory was predicted using a premeasured gradient impulse response function.[9] The multiecho reconstruction was performed prior to NUFFT[10] (30 Hz Lorentzian filtered; reconstructed resolution 5.62 × 5.62 mm²; nominal 128 × 128 matrix). n = 1 control and infarcted animals were scanned; other details as above.
Results
Myocardial infarction was confirmed via CINE. The $[1,4\text{-}^{13}\text{C}_2]$malate:$[1,4\text{-}^{13}\text{C}_2]$fumarate ratio visible in the in vivo infarcted heart was 0.21 ± 0.03 one day post infarction, dropping to 0.077 ± 0.005 seven days post infarction compared to 0.0025 ± 0.001 in controls, corresponding to an ~ 82-fold increase in visible malate signal post-infarction (Figs. 2&3; Cohen’s $d = 5.01$). Very little $[1,4\text{-}^{13}\text{C}_2]$malate was observed in control animals, potentially indicating the negligible rate of transport of the probe within the lifetime of the hyperpolarized experiment. As shown in Fig. 4, it was possible to resolve fumarate perfusing the heart, and malate in the region of the infarction in the basal LV in the infarcted animal shown; production was not visible in controls. This region of necrosis was smaller than the corresponding wall motion abnormality in CINE, and smaller than the detected region of enhancement by late gadolinium.

Discussion
Malate production in the infarcted heart appears promising as a sensitive in vivo probe of necrosis following MI. We have shown that the technique could offer the first non-invasive measurement of necrotic cell death in vivo. The region of malate signal detected in the heart following MI appears distinct from the region of late gadolinium enhancement, which is consistent with gadolinium revealing aberrant vasculature and regions of fibrosis rather than necrosis directly. Hyperpolarized $[1,4\text{-}^{13}\text{C}_2]$fumarate is therefore suited to studying the decline into heart failure following MI. In contrast to the use of blood-borne biomarkers such as troponin to detect necrosis, hyperpolarized $[1,4\text{-}^{13}\text{C}_2]$fumarate can be spatially localised, and the rapid timescale of hyperpolarized experiments is uniquely suited to acute MR investigations.

Conclusions and Future Work
Hyperpolarized $[1,4\text{-}^{13}\text{C}_2]$fumarate can be used to specifically identify cardiomyocyte necrosis both acutely following myocardial infarction and up to one week later. Future work will investigate its co-localization to the region of myocardial injury, and investigate the specificity and sensitivity of the technique in population studies compared to troponin.

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References

Figures
Fig. 1: The designed hybrid excitation pulse sequence (A), together with the nodal excitation plot (B), which excites both malate resonances with approximately a 20° flip angle and fumarate at 4°.

Fig. 2: Example spectra acquired from (i) control animals or those 1-day (ii) or 7-days (iii) post infarction. The visible hyperpolarized (HP) malate signal is substantially increased one day post infarction compared to control animals when normalised to either a thermally polarized (TP) $^{13}$C urea reference or the injected fumarate peak. The visible malate signal reduces over time, reflecting the progression of the disease.

Fig. 3: The timecourse of disease is clear across the three groups of animals scanned, with a significant difference in malate production observed between all three groups ($\ast$ denotes Student’s $p < 0.05$).
Fig. 4: Visible reconstructed [1,4-$^{13}$C$_2$]fumarate and [1,4-$^{13}$C$_2$]malate images from the infarcted heart. No malate signal was resolvable in healthy animals. The detected malate signal co-localised to the region of the infarct as determined by CINE and late gadolinium (arrow).