Magnetic resonance fingerprinting of the pancreas at 1.5 T and 3.0 T

Eva M. Serrao1,2,3, Dimitri A. Kessler1,2, Bruno Carmo1,2, Lucian Beer1,2, Kevin M. Brindle3, Guido Buonincontri4, Ferdia A. Gallagher1,2,3, Fiona J. Gilbert1,2,3, Edmund Godfrey4, Martin J. Graves1,2, Mary A. McLean1,2,3, Evis Sala1,2,3, Rolf F. Schulte5 & Joshua D. Kaggie1,2

Magnetic resonance imaging of the pancreas is increasingly used as an important diagnostic modality for characterisation of pancreatic lesions. Pancreatic MRI protocols are mostly qualitative due to time constraints and motion sensitivity. MR Fingerprinting is an innovative acquisition technique that provides qualitative data and quantitative parameter maps from a single free-breathing acquisition with the potential to reduce exam times. This work investigates the feasibility of MRF parameter mapping for pancreatic imaging in the presence of free-breathing exam. Sixteen healthy participants were prospectively imaged using MRF framework. Regions-of-interest were drawn in multiple solid organs including the pancreas and T1 and T2 values determined. MRF T1 and T2 mapping was performed successfully in all participants (acquisition time: 2.4–3.6 min). Mean pancreatic T1 values were 37–43% lower than those of the muscle, spleen, and kidney at both 1.5 and 3.0 T. For these organs, the mean pancreatic T2 values were nearly 40% at 1.5 T and < 12% at 3.0 T. The feasibility of MRF at 1.5 T and 3 T was demonstrated in the pancreas. By enabling fast and free-breathing quantitation, MRF has the potential to add value during the clinical characterisation and grading of pathological conditions, such as pancreatitis or cancer.

Magnetic resonance imaging (MRI) of the pancreas is increasingly used as a major diagnostic modality for characterisation of pancreatic lesions, given its superior soft tissue contrast and increased sensitivity for detection and characterisation of smaller pancreatic masses1,2. However, the wider use of MRI remains hampered by long examination times, which limits the types of acquisition to the minimum required for basic diagnosis. Contrast-enhanced CT is routinely used in the context of pancreatic disease given its wide availability and acquisition time. However, radiation exposure and iodine allergy are its major risks. Endoscopic ultrasonography (EUS) is the most sensible imaging method for the pancreas3,4, but it is invasive, operator dependent and associated with complications.

T1 and T2 relaxation times are valuable quantitative parameters to characterize different tissues, particularly in the assessment of myocardial and liver diseases5–9. Thus far only some studies have reported on the value of quantitative MRI on the pancreas. In normal pancreas, T1 values have been reported to reflect the amount of acinar protein, rough endoplasmic reticulum and fat infiltration10–13. T1 maps were found to aid determining the presence and severity of acinar cell loss in the diagnosis and classification of chronic pancreatitis10,11,14. Multiparametric MRI comprising T1, T2, and ADC mapping was also shown useful in discriminating different pancreatic processes15. Despite the potential of quantitative multiparametric MRI, pancreatic MRI protocols are still mostly qualitative, with clinical assessments involving a trained reader to create a subjective evaluation based on T1- and T2-weighted images. This subjective evaluation is highly parameter dependent, which reduces the ability for analysis to be translated across centres. Quantitative analysis is mostly undertaken in a research context, often limited to conventional methods, due to the significant scan time required and technical limitations such as field inhomogeneities and patient motion9,16.

Several quantitative methods have been described for fast abdominal imaging17–19. MR fingerprinting (MRF) is an innovative technique that provides qualitative and quantitative data from a single exam19. As a T1 and T2 mapping method, MRF has demonstrated itself as a fast, repeatable within a system and reproducible across centres20. MRF has been shown to be quite insensitive to motion due to incoherent sampling resulting from the golden angle rotations, which enable pattern matching if a voxel is static for sufficient frames21. MRF-derived multiparametric maps have shown diagnostic utility in the brain22–24 and abdomen25–27. In tumours,
MRF measurements have shown T1 values in lesions that are nearly double those of normal-appearing tissue in the prostate, liver and brain, and demonstrated T2 differences between low and high grade tumours as great as 70%24–27. The application of MRF in the pancreas, and in the abdomen in general, offers a wide range of clinical and research opportunities by accelerating acquisition of quantitative parameter maps.

The challenges associated with MRF are similar to those for any quantitative MRI technique: motion, spatial resolution, and B0 and B1 field non-uniformity. MRF may also be sensitive to magnetization transfer, partial volume, and slice profile effects21,28–32. Spiral sampling use in MRF is advantageous in that it may reduce motion sensitivity by oversampling the centre of k-space33 and matching a temporal pattern with MRF reconstruction.

This work investigates the feasibility of MRF for pancreatic imaging in the presence of free-breathing motion for the first time. Pancreatic multiparametric MRI has shown potential clinical impact in the assessment of pancreatic disease. We predict that MRF acquisition might be of particular value in pancreatic cancer (PCa) patients, who are often frail and therefore less compliant, allowing for improved quantitative clinical assessment to be performed. Most MRF investigations published to date have been performed at 3.0 T; however, the use of MRF at 1.5 T would increase its clinical potential given the wider availability and clinical indications for these systems. Here we compare the performance of MRF in the abdomen at both 1.5 T and 3.0 T in healthy subjects.

Results

Feasibility of MRF of the pancreas at 1.5 T and 3 T. MRF T1 and T2 maps were obtained for each healthy volunteer. The mean age for the 16 participants was 33 years, of which 6 (40%) were female. Sixteen participants (100%) underwent MRI at 3.0 T and 12 (80%) underwent MRI at 1.5 T. The volunteers had normal appearances of the liver, pancreas, kidneys and spleen as assessed by a radiologist. The MRF acquisition time was 146–215 s (2.4–3.6 min) per subject. Dictionary simulation normally required approximately 45 min for reconstruction with parallelized code. The reconstruction of each channel, slice, and frame was the largest time-consuming step prior to compression of the dataset with singular value decomposition (SVD) factorization. After SVD compression, the T1 and T2 maps were determined by finding the maximum inner product between the simulated signal and the acquired images. Product pattern matching took a few seconds. The abdominal images required 3.5 h for non-Cartesian reconstruction and matching due to the large number of reconstructed frames and coil channels.

Quantitative and qualitative interpretation of the MRF maps. Axial and coronal images were acquired at 3.0 T in the same subject. Coronal acquisition was found to be more optimal to image the upper abdomen (pancreas), avoiding most of the artefacts encountered with axial imaging (Fig. 1). These artefacts, created primarily due to motion but also undersampling, caused underestimation of the T1 means for each
single subject, in example the T1 mean value of the whole kidney reduced from 2071 ± 428 ms (coronal) to 1045 ± 369 ms (axial). In the pancreas the T1 means was up to 100 ms higher in the coronal plane.

T2-weighted single-shot-fast-spin-echo sequence (SSFSE) images and MRF T1 maps are shown in Fig. 2. These images highlight the good anatomic detail obtained by MRF maps when compared with the conventional SSFSE image. The liver and pancreas (arrowed) have similar intensities in the T1 maps, while fat, with a much shorter T1, appears dark and muscle, with a longer T1 appears bright. SSFSE: single-shot-fast-spin-echo sequence, MRF: magnetic resonance fingerprinting.

Figure 2. (a, c) T2-weighted SSFSE images and (b, d) MRF-derived T1 maps of the abdomen at (a, b) 1.5 T and (c, d) 3.0 T. This demonstrates the good anatomic detail and homogeneous signal throughout the pancreatic gland, obtained by MRF maps when compared with the conventional SSFSE image. The liver and pancreas (arrowed) have similar intensities in the T1 maps, while fat, with a much shorter T1, appears dark and muscle, with a longer T1 appears bright. SSFSE: single-shot-fast-spin-echo sequence, MRF: magnetic resonance fingerprinting.

Tissue-specific T1 and T2 values, at 1.5 T and 3 T, are reported in Tables 1 and 2 (pancreas and non-pancreas, respectively) and Fig. 5. The pancreas (head, body, and tail) showed relatively homogenous appearances and T1 and T2 values, which are listed in Table 1. No significant differences in T1 and T2 values were found between the pancreatic head, body, and tail (p > 0.05).

As expected, T1 relaxation times for all organs were significantly longer at 3 T (1.5 T 1.02 ± 0.3 s vs. 3.0 T 1.26 ± 0.38 s, p = 0.0001). Similar T1 and T2 trends between organs were observed at both field strengths, although the trends were more pronounced for both T1 and T2 at 1.5 T and for T1 at 3.0 T. The liver and pancreas could not be differentiated based on the T1 or T2 relaxation values at both field strengths.

The pancreatic T1 values were 30–50% lower than in muscle, spleen, and kidney at both 1.5 and 3.0 T. Large standard deviations were found for the spleen, muscle and kidney. However, in the case of the kidney the drawn ROI included both medulla and cortex, which have different relaxation times.

Phantom results. The phantom results are shown in Fig. 6, demonstrating the results of several T1 and T2 mapping methods.

For T1 mapping, these show that MRF agrees with fast spin echo with inversion recovery (FSE-IR) values, which is considered close to gold standard. The 2D variable flip angle (VFA) method vastly underestimated T1, whereas the 3D VFA method overestimated T1. The modified Look-Locker inversion recovery (MOLLI) approach underestimated T1 mildly, when compared with FSE-IR above 500 ms; below 500 ms, the MOLLI-T1
values plateaued. These biases were similar between both 1.5 T and 3.0 T results. As a quantitative metric of agreement, the sum across all 28 vials of the absolute differences, and divided by 28 to account for additional samples, between the $T_1$ methods and the FSE-IR results were: for 3.0 T, 0.16 s for MRF, 0.18 s for MOLLI; 0.27 s for 3D VFA; and 0.50 s for 2D VFA; and for 1.5 T, 0.14 s for MRF, 0.24 s for MOLLI; 0.25 s for 3D VFA; and 0.50 s for 2D VFA. We note that this measurement is biased due to the higher numbers of low-value $T_1$ vials, which increases their importance or weighting in this calculation; when low-value $T_1$ vials (< 100 ms) were excluded, this mean of absolute differences between MRF and FSE-IR was 0.05 s at 3.0 T and 0.03 s at 1.5 T, while other methods were greater than 0.15 s.

For $T_2$ mapping, both multi spin echo (MSE) and MRF agreed up until 300 ms, at which point the values diverged. This is accounted for by not obtaining a sufficient number of lengthy TEs during the MSE acquisition.

Discussion
This study demonstrates the feasibility of free-breathing, non-gated MRF in the pancreas at 1.5 and 3 T within a clinically reasonable acquisition period of 2.4–3.6 min. The MRF framework allows qualitative and quantitative data to be acquired simultaneously, allowing ready comparison between longitudinal time points and against population-derived norms, as well as giving improved imaging repeatability and more meaningful interpretation of intensity changes. The total acquisition time was very low when compared with other reported single parameter acquisitions (non-MRF based), which include quantitative $T_2$ measurements of the pancreas at 1.5 T: 8 min\(^{16}\) and 3 T: 2 min 50 s\(^{9}\). Similar to Wang\(^{35}\), $T_1$ maps with MRF were obtained in 10 s/slice, although we did

Figure 3. MRF-derived maps of (a) relative proton density (rPD), (b) $T_1$, and (c) $T_2$ of three consecutive slices (posterior to anterior) within the abdomen at 1.5 T. These images show the ability of MRF to obtain multiple slices through the abdomen with reasonable anatomical detail and low motion artefact. The pancreas (arrowed) has homogeneous signal throughout in both $T_1$ and $T_2$ maps, and is distinguishable due to fat/water boundaries at its periphery.
not perform breath-holding and obtained images at nearly half the voxel volume. Furthermore, the repeatable\textsuperscript{20} quantitative nature of MRF data has the potential to improve comparability between centres.

Despite the overall lack of motion sensitivity in MRF\textsuperscript{21}, we found coronal acquisition to be preferred over axial. The axial artefacts were found to be due to motion rather than FOV limitation. Prior MRF studies\textsuperscript{25} do not report such limitations in axial images, likely because different MRF parameters were applied, leading to greater SNR at the cost of increased acquisition time. The respiration artefacts were limited in the coronal plane as the motion remained in-plane through all excitations. In the axial plane motion occurs in the slice direction, leading to underexcitation of through-plane voxels and consequently incorrect T1 and T2 values. Motion occurred occasionally in coronal acquisitions resulting in inaccurate T1 and T2 values. We found that air in the gut lead to higher field non-uniformities in T2 maps.

Here we report MRF-derived quantitative differences between normal tissue types, their T1 and T2 relaxation values and the relationship of these values with field strength. Similar T1 and T2 trends between organs were observed at both field strengths, although the trends were more pronounced for both T1 and T2 at 1.5 T and for T1 at 3.0 T. As per previous literature\textsuperscript{34}, there was no significant difference (p > 0.05) in the T1 or T2 relaxation values of liver and pancreas, at both field strengths. The pancreas had lower T1 and T2 values than muscle, spleen, and kidneys at both field strengths. The mean T2 values of the pancreas were nearly 40% lower than those of muscle, spleen and kidney at 1.5 T. This suggests that MRF-derived T2 maps at this field strength can be used to discern these organs more easily than at 3 T, where the mean T2 values varied by < 12%.

Pancreatic processes, including cancer, have long T1 values (Table 2), which can be challenging to map with most techniques, as they result in lower signal recovery between pulses and require longer recovery periods for

Figure 4. (a) Relative proton density (rPD), (b) T1 maps, and (c) T2 maps of three consecutive slices (posterior to anterior) within the abdomen at 3.0 T. These images show the ability of MRF to obtain multiple slices through the abdomen with reasonable anatomical detail and low motion artefact. Again, homogeneous signal was shown throughout the pancreas (arrowed). These T2 maps result in larger spiral artefacts that are not present on the T1 maps or at 1.5 T.
full signal relaxation. However, MRF has been reported to measure long \( T_1 \) values accurately\(^{20,26} \), while remaining time-efficient. This study shows the ability of MRF to accurately acquire \( T_1 \) and \( T_2 \) maps of the normal pancreas, despite its relatively long \( T_1 \) and \( T_2 \) values. The signal homogeneity throughout the normal pancreas in both \( T_1 \) and \( T_2 \) maps at 1.5 and 3.0 T was a key observation, as it strengthens the use of MRF in the context of pancreatic disease. This will potentially allow depiction of regional variability/heterogeneity, and definition of boundaries between pancreatic processes and normal pancreatic tissue within the same patient, which would have clear clinical impact, particularly in the context of PCa. Clinical overlap between PCa and chronic pancreatitis (CP) is well recognised, as CP increases the risk of PCa and often coexist with PCa\(^{33,36} \). We envisage, that distinguishing the two processes by MRF might prove challenging, but further studies with histopathological correlation will be needed.

Despite the higher \( T_1 \) values at 3.0 T (1010–1041 ms) obtained in this study, these remained 300–700 ms lower than those measured in patients with pancreatic disease using MOLLI (1324 ms for CP; 1675 ms for pancreatic ductal adenocarcinoma)\(^{35} \), suggesting that MRF would still be able to distinguish normal pancreatic tissue from diseased pancreas. However, we envisage that if MRF in coronal plane was to be used, then disease pancreas would also have proportionally higher values. Ascites is often present in patients with advanced PCa.

**Table 1.** Mean and standard deviation of \( \text{T}_1 \) and \( \text{T}_2 \) values for each pancreatic region. The references are: \( \gamma^{34} \), \( \delta^{16} \), \( \phi^{35} \), \( \&^{15} \).

| Region             | \( \text{T}_1 \) (ms) | \( \text{T}_2 \) (ms) |
|--------------------|-----------------|-----------------|
| Pancreas head      | 798 (67)        | 65 (10)         |
| Pancreas body      | 799 (72)        | 59 (16)         |
| Pancreas tail      | 803 (53)        | 68 (14)         |
| **Average**        | 584 (14)        | 1029 (65)       |

\( ^{1} \) \( \gamma^{14} \), \( ^{4} \) \( \gamma^{8} \), \( ^{5} \) \( \delta^{9} \), \( ^{6} \) \( \phi^{10} \), \( ^{7} \) \( \&^{15} \), \( ^{8} \) \( \delta^{16} \), \( ^{9} \) \( \phi^{35} \), \( ^{10} \) \( \&^{15} \).
The presence of large volume of ascites impacts on image quality, as it creates B1 field non-uniformity. However, previous studies from our group have shown that high quality maps could be obtained in patients with large ascites.

The MRF-T1 values at 1.5 T and 3.0 T were shown to be 27% (584 ms) and 15% (865–884 ms) higher than in the literature, where MOLLI was used for in vivo measurements, respectively\cite{10,14,34,35}. However, MOLLI can be affected by variable heart rates, incomplete tissue recovery between inversion pulses, and adiabatic inversion inefficiencies resulting in an underestimation of T1 by as much as 25% at 3.0 T\cite{37} and 17% at 1.5 T\cite{38}. Similarly to our results, T1 miscalculation has been reported with cardiac MRF, where the MRF-T1 values were 173 ms higher than MOLLI-T1 values\cite{36} and 97–189 ms lower than SASHA-T1 values\cite{39,40}. Paradoxically, our phantom work demonstrated lower MRF-T1 values than MOLLI-T1 values, indicating the complexity and multifactorial

| Region  | 1.5 T | 3.0 T |
|---------|-------|-------|
| Liver   | 774 (62) | 586 (39) |
| Muscle  | 1253 (92) | 856 (61) |
| Spleen  | 1420 (105) | 1057 (42) |
| Kidney  | 1503 (200) | 1189 (58) |

Table 2. Mean and standard deviation of T1 and T2 values of non-pancreatic abdominal regions. All measured regions had T1 values 8–28% longer and T2 values 36–80% shorter at 3.0 T when compared with 1.5 T. The reference for γ is 34 and ‡ is 25.

![Figure 6. T1 and T2 phantom values at both 1.5 T and 3.0 T, using conventional techniques and MRF. T1 values between MRF and FSE IR had low absolute mean differences (<0.05 s for all vials above 0.1 s). There were four vials where the 2D VFA, 3D VFA, and 2D MRF all failed, which occurred at the centre of the T2-layer of the NIST phantom where the T1 values were <30 ms. The MSE flattened at longer T1 values. MOLLI-T1 and 3D VFA was higher than both MRF-T1 and FSE-IR-T1.](image-url)
nature behind the in vivo measurements. Our MRF-T₂ results were also ~200 ms longer than those reported by prior MRF studies¹⁵,²⁵, which more closely matches MOLLI values. We found that this difference was likely due to a combination of technical and biological factors, as we included different MRF parameters, acquisition plane and a younger cohort. Coronal acquisition was found to increase T₁ values when compared with axial acquisition (used in the other studies). The significantly younger mean age of our cohort might have contributed to a lesser extent to our results. The MRF T₂ relaxation times, at both field strengths, were also in the higher range, which could be due to factors inherent to the sequence⁷.

In this study, we also found that organs lying at the image periphery, such as muscle, kidney and spleen, showed a wide T₁ and T₂ SD values and higher T₁ mean differences between acquisition plane, independently of the field strength, likely due to MRF inherent k-space undersampling at the edges causing motion-like artefacts and poor dictionary matching. These artefacts could be reduced by increasing voxel size or the number of frames such that the SNR is increased at the cost of increased time acquisition. Also, the T₁ and T₂ values showed multimodal distribution for both field strengths and tissue type, which was partially caused by the few number of participants (n = 16). The distributions for each organ did not simply shift between field strengths, but resulted in non-trivial transformations, likely due to slightly different analysed anatomic locations as an effect of patient position/respiration. Despite these limitations, MRF is one of few methods that can obtain free-breathing T₁ and T₂ parameter maps within reasonable acquisition times.

This work was also challenging due to computational limitations involving the high dimensionality of the acquired and simulated datasets, which limited either acquired raw data or matching of transient state simulation parameters. The approximately 4 min abdominal MRF scan used approximately 4 gigabytes of disk space due to near continuous data acquisition combined with the large number of coil channels. The original MRF paper⁴⁹ used 32 coil channels, a matrix size of 128 × 128, a single slice, and 1000 frames, whereas our work uses near 20 slices with twice the matrix sizes. Reconstruction prior to dictionary matching required memory reduction steps, such as coil combination and SVD compression. During dictionary matching where the inner product was calculated between the simulated dictionary and compressed acquisition data, the maximum amount of RAM used was 350 gigabytes while using 44 threads (Xeon Gold 6152). When B₁ or B₀ values were simulated, the T₁ and T₂ maps appeared much noisier, and therefore these were not performed.

This proof-of-principle study has shown the feasibility of using free-breathing, non-gated coronal MRF for fast imaging and quantification of relaxation parameters in the normal pancreas. We envisage that the MRF framework will be of great value in patients with PCa, who are usually frail and with limited tolerance to long examinations or breath-hold MRI measurements. The MRF technique might also prove useful in characterising and grading pathological conditions such as CP, given its ability to acquire simultaneous mapping of T₁ and T₂ as well as qualitative images. Furthermore, the demonstrated feasibility of MRF in the abdomen at 1.5 T could significantly impact on the clinical potential of MRF as an imaging tool, as 1.5 T remains the most widely used field strength worldwide.

Materials and methods
Sixteen healthy volunteers were imaged in the supine position with free-breathing MRF using a 32-channel abdominal array on a 3.0 T MRI system, after informed consent, with twelve of the sixteen also imaged on a 1.5 T MRI system (MR750 and MR450, GE Healthcare, Waukesha, WI, USA, respectively). Fasting was not requested to the volunteers.

The present study protocol was reviewed and approved by the Hertfordshire Research Ethics Committee (REC ref 08/H0311/117, IRAS 161555, REC approval on 12 Sept 2008). The present study was performed in accordance with relevant guidelines and regulations. To protect the individuals’ privacy, the patient’s exam information was pseudo-anonymised by replacing personal identifiers with pseudonyms. All work was carried out in accordance with relevant guidelines and regulations.

Phantom protocol. MRF data were acquired with an inversion-prepared 2D steady-state-free-precession (SSFP) MRF sequence (1, 2). The acquisition consisted of 979 undersampled interleaved spirals with 656 points per spiral, and with sequential spirals rotated by the golden-angle (Fig. 7a). The maximum gradient strength per spiral was 28 mT/m and the maximum slew rate was 108 T/m/s. The imaging parameters were: field-of-view (FOV) = 260 × 260 mm², matrix = 256 × 256, slices = 3, slice thickness = 3.0 mm, spacing 1.0 mm, sampling bandwidth = ±250 kHz, slice dephasing = 8°, echo time (TE) = 2.5 ms, repetition time (TR) = 10 ms, acquisition time = 9.79 s/slice. The flip angle lists matched those in Jiang et al.⁴¹ (Fig. 7b). A static TR was used as the random TRs listed in Jiang et al.⁴¹ gave an unpleasant auditory pitch.

For VFA-T₁ mapping, both 2D and 3D data were acquired with a fast spoiled gradient echo (FSPGR) method using flip angles of 2, 5, 8, 12, 15, 18, 22, and 26°. The 2D data matched the FOV, matrix, and slices as the MRF acquisition. For FSE-IR-T₁ mapping, data were acquired with inversion times (TIs) of 50, 100, 200, 400, 800, 1600, and 2400 ms. T₁ data for all non-MRF techniques were fit using non-linear fitting of the signal equations.

Multi spin echo (MSE) data for T₂ estimations were acquired with TEs = 8.1, 16.3, 24.4, 32.6, 40.7, 58.9, 57.0, 65.2 ms. MSE data were fit with a log linear least squares algorithm.

In vivo protocol. MRF data were acquired with the same parameters as in the phantom. Coronal and axial images were acquired at 3.0 T, but only coronal images were acquired at 1.5 T due to fewer respiratory artefacts than axial images.

Images were also obtained with a coronal single-shot-fast-spin-echo sequence at 3.0 T during a 33 s breath-hold with TR = 1132 ms, TE = 80 ms, matrix = 448 × 224, field-of-view = 360 × 324 mm², slice thickness = 6 mm,
slices = 24, bandwidth = ± 83.33 kHz, coil acceleration factor = 2. At 1.5 T, with TR = 1102, TE = 62 ms, matrix = 320 × 224, FOV = 460 × 460 mm², slice thickness = 5 mm, slices = 26, and bandwidth = ± 83.33 kHz.

**MRF Image Reconstruction.** Each under-sampled spiral was reconstructed to give 979 under-sampled images per slice. The spiral k-space was regridded and interpolated to a Cartesian k-space before a Fast Fourier Transfer (FFT), and used a three frame sliding window. The images were reconstructed with 48 parallel CPUs and used 400 gigabytes of RAM. After reconstruction, each coil channel was combined using adaptive coil combination based on weights determined from the average of the time frames. The undersampled images were reduced from 979 to 16 images using the SVD decomposition weights determined during dictionary compression.

**MRF dictionary simulation.** Dictionary simulations of the signal evolution in a steady-state-free precession acquisition scheme were performed using the extended phase graph formalism. The slice profile was also included. The ranges and incremental (step-size) changes of the $T_1$ and $T_2$ values that were simulated in the dictionary were $T_1 = [0.01:0.005:1;1:0.04:6]$ seconds ([minimum: step-size: maximum]), and for $T_2 = [0.005:0.001:0.1; 0.1:0.01:4; 4:0.04:6]$ seconds (where the semi-colons indicate concatenated lists). The dictionary size was compressed to 16 singular vectors (rank) with SVD to reduce the size for long term storage and faster dictionary matching.

**MRF pattern matching.** MRF uses a pattern recognition algorithm to identify the $T_1$ and $T_2$ tissue properties in each voxel. The $T_1$ and $T_2$ maps from MRF were obtained by inner product pattern matching of the dictionary, which is a signal look-up table based on simulations with different $T_1$ and $T_2$ times, with the best match to the acquired reconstructed data.

The inner products between the normalized measured signal evolution of each voxel and each normalised dictionary entry are calculated. The dictionary entry returning the maximum value for the inner product is taken as the best representation of the acquired signal evolution. The respective $T_1$ and $T_2$ values are consequently assigned to the voxel. The rPD (relative proton density) is calculated as the scaling factor used to match the dictionary simulation with the measured signal evolution.

**Region-of-interest selection.** Pancreatic (head, body and tail), liver (right hepatic lobe avoiding the inclusion of vessels), kidney (most cases lower pole, including cortex and medulla), spleen and muscle (right psoas muscle) sub-regions (regions of interest) were identified and drawn manually in both $T_1$ and $T_2$ MRF maps, for each subject and imaging exam, via tracing with a computer mouse by a trained reader using custom in-house software.

**Statistical analysis.** Data were analyzed using GraphPad Prism v6 (GraphPad Software, San Diego, USA). Data were reported as mean ± SD, unless stated otherwise. Statistical significance was tested with Prism using unpaired Student’s $t$-test and ANOVA tests to compare quantitative parameters between groups. The results were considered to be significant when $p < 0.05$.

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**Figure 7.** (a) The acquisition consisted of undersampled spirals that were rotated by the gold-angle after each TR. The first nine spirals are shown here. (b) One spiral (or frame) was acquired per TR, with the flip angle varied per TR as shown here.
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**Author contributions**

E.M.S. and J.K. conceived and designed the study. E.M.S., J.K., D.A.K., M.M. and B.C. performed experiments and collected data. G.B., M.M., M.J.G. and R.F.S. provided technical support. E.G. reviewed the images clinically. E.M.S. and J.K. analysed data. E.M.S., J.K. wrote and K.M.B., F.A.G., L.B., F.J.G. and E.S. provided research support. All authors reviewed and approved the manuscript.

**Competing interests**

The authors declare no competing interests.

**Additional information**

**Correspondence** and requests for materials should be addressed to J.D.K.

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