High-Resolution Self-Gated Dynamic Abdominal MRI Using Manifold Alignment

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Abstract— We present a novel retrospective self-gating method based on manifold alignment (MA), which enables reconstruction of free-breathing, high spatial and temporal resolution abdominal MRI sequences. Based on a radial golden-angle (RGA) acquisition trajectory, our method enables a multi-dimensional self-gating signal to be extracted from the k-space data for more accurate motion representation. The k-space radial profiles are evenly divided into a number of overlapping groups based on their radial angles. MA is then used to simultaneously learn and align the low dimensional manifolds of all groups, and embed them into a common manifold. In the manifold, k-space profiles that represent similar respiratory positions are close to each other. Image reconstruction is performed by combining radial profiles with evenly distributed angles that are close in the manifold. Our method was evaluated on both 2D and 3D synthetic and in vivo datasets. On the synthetic datasets, our method achieved high correlation with the ground truth in terms of image intensity and virtual navigator values. Using the in vivo data, compared to a state-of-the-art approach based on centre of k-space gating, our method was able to make use of much richer profile data for self-gating, resulting in statistically significantly better quantitative measurements in terms of image sharpness and image gradient entropy.

Index Terms— manifold alignment, MRI self-gating, MRI reconstruction, respiratory motion.

I. INTRODUCTION

DYNAMIC magnetic resonance imaging (MRI) involves imaging a region of interest with high temporal resolution, and is useful in many applications in which knowledge of motion is of interest. For instance, dynamic MRI can help in assessing coronary artery disease [1], motion correction of simultaneously acquired PET data [2, 3] or for studying the nature of respiratory motion [13]. These last two applications require a long MRI acquisition (e.g. up to 10 minutes in [3]) to be performed. However, the acquisition speed of MRI prevents sufficient data from being acquired quickly enough to reconstruct high-resolution fully sampled images (in both 2D and 3D). This problem can be tackled by using under-sampled reconstruction schemes such as compressed sensing (CS) [4]. CS typically involves an iterative optimisation process, which can be time-consuming for large amounts of dynamic MRI data. Furthermore, motion may still occur during the period required to acquire the under-sampled k-space data, especially in 3D image acquisition. To tackle this problem, a gating approach can be used. This involves the combination of corresponding k-space data that were acquired at different times but similar motion states. Gating typically relies upon a gating signal to establish these correspondences. A range of different gating signals have been proposed, such as external sensing based techniques [5, 6], a pencil-beam navigator [7] and self-gating methods [8, 9]. Examples of external sensing based methods include the use of optical tracking devices [10] or respiratory bellows [6]. Such techniques often involve additional setup time, may interfere with the magnetic resonance signal or may suffer from limited field of view or line-of-sight problems (e.g. optical tracking).

In contrast, self-gating methods address these problems by performing the gating using the acquired data themselves. A common approach is to use the magnitude of the centre or central line of k-space which is acquired continuously during the acquisition as the self-gating signal [11, 12].

Most external sensing-based methods, as well as pencil-beam navigator and current self-gating techniques, only estimate a simple 1D gating signal (typically in the head foot direction). However, organ motion is a complex phenomenon (e.g. diaphragm contraction and rib cage motion in respiratory motion), so the use of such simple signals for gating limits the quality of the reconstructed images. von Siebenthal et al. [13] developed a 4D MR imaging method based on stacking of dynamic 2D images using internal image-based sorting, which achieved better image quality than using the 1D gating signal.

Our proposed method, which is based on manifold alignment (MA), is a self-gating method which enables a multi-dimensional signal to be used for gating. Similar to the centre of k-space gating (CKG) method, the intuition behind our technique is that respiratory motion is pseudo-repetitive in nature, and can therefore be represented by a small number of motion variables. MA is used to uncover these underlying variables. A number of previous works have reported the use of manifold learning or MA for self-gating in MRI [2, 14-20]. Usman et al. [20] applied Laplacian Eigenmaps to estimate respiratory motion from the central intersection region of a number of consecutive k-space profiles using a radial golden-angle acquisition (RGA) [21]. Similarly, Bhatia et al. [15] used Laplacian Eigenmaps to estimate cardiac motion from repetitively sampled central k-space lines using a...
Cartesian acquisition. Recently, Poddar et al. [18] used Laplacian Eigenmaps to estimate both cardiac and respiratory motions from repetitively sampled radial trajectories. All of these methods were only applied in 2D acquisition, and their extension to 3D acquisition is not trivial as the repetitive sampling strategy reduces the scanning efficiency. In [2], we proposed a method for MRI self-gating using MA of 2D reconstructed slices acquired at different anatomical positions. The technique worked by aligning the manifolds of manifold across slices in a group-wise fashion. Correspondences were established in the manifold and corresponding slices were stacked into volumes. In [14] the technique was extended by embedding all 2D slices simultaneously. This was achieved by avoiding inter-slice data comparisons by representing each slice using a novel feature descriptor based on graph theory and random walks [22]. However, the methods in [2, 14] are image-based self-gating methods that assumed there was no motion within each of the fully sampled 2D image acquisitions (~300 ms per image), which may not be true for fast motion. The use of images for self-gating also limits the achievable temporal resolution to the time taken to acquire enough data to reconstruct an entire image. To improve this technique, we have recently proposed a MA framework that is directly applied to k-space profiles [16] to achieve dynamic volume reconstruction based on a multi-slice 2D acquisition. It simultaneously embeds k-space profiles that are acquired at different slices and temporal positions into a common manifold. 3D dynamic sequences can be reconstructed with an improved temporal resolution of ~45 ms. The above multi-slice 2D acquisition methods [2, 14, 16] assume that the underlying motions of all slices are similar which is true for small field-of-view (FOV) imaging (e.g. liver and heart). However for a large FOV like the whole thorax/abdomen, it is not guaranteed that slice correspondences can be accurately established using MA.

In this paper, we adapt the MA method in [14] and propose a novel framework based on k-space MA for high spatial and temporal resolution MRI reconstruction, which is suitable for wide FOV imaging. The focus of this paper is on respiratory motion, although we believe that the technique may also be applicable to cardiac motion. Some of our prior work has been presented in [17]. We describe here the full details of the technique with some improvements, and report comprehensive evaluation results. The main novelties and contributions of this work are summarised as follows: (1) In contrast to [15, 20] and the CKG method, instead of using a 1D gating signal, the proposed method performs gating in a low dimensional manifold. The use of a multidimensional gating signal and reconstruction of a high temporal resolution sequence allows better representation of the intra-cycle and inter-cycle motion variations than the CKG reconstruction of a limited number of motion states. (2) In contrast to [14], we perform MA directly on k-space data rather than the reconstructed image. This enables our technique to achieve higher temporal resolution of the reconstructed volumes, as the fully sampled 2D image requires ~300 ms per image, and the acquisition of a k-space profile only takes ~3 ms. (3) Additional to [17], we use overlapping k-space groups and an additional cost function term to achieve a more reliable MA. (4) In contrast to [2, 14, 16], our method works using a 3D acquisition rather than a multi-slice 2D acquisition, thus avoiding the assumption of a similar underlying motion across all slice positions. (5) The proposed method allows reconstruction of images at as many respiratory positions as the number k-space profiles (2D) or the number of stacks of k-space profiles (3D) acquired. (6) Most importantly, we demonstrate that our method works for both 2D and 3D acquisitions.

**II. Methodology**

Fig. 1 provides an overview of the proposed method, which consists of k-space data acquisition, MA, and image reconstruction. During data acquisition, we acquire k-space profile data continuously under free-breathing with a RGA

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**Fig. 1. Overview of the MA method for self-gating dynamic MRI with the following steps. (a) K-space data is acquired using RGA trajectory. (b) All k-space radial profiles are assigned into G overlapping groups according to their profile angles. The profiles are embedded in a low dimensional manifold using MA. (c) Image reconstruction at each profile position by combining nearby profiles with Euclidean distance weighted contributions.**

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**Respiratory cycles under free breathing**

(a) K-space radial golden angle acquisition

(b) Manifold alignment of K-space data

(c) Image reconstruction at each profile position by combining nearby profiles with Euclidean distance weighted contributions.
trajectory. The temporal series of k-space lines (profiles) are then assigned to overlapping groups based only on their radial angle. Therefore each k-space group, whilst only containing a limited range of angles in k-space, will contain data acquired at many different time points throughout the entire range of motion states. As such, the data in a given k-space group lie on a low dimensional manifold which captures the range of motion states encountered during data acquisition. Rather than learn the manifold for each k-space profile group individually, MA is used to simultaneously embed all k-space groups within a common low-dimensional manifold. In the manifold, k-space profiles that were acquired at similar motion states (respiratory positions) are close to each other. For each acquired k-space profile, its manifold coordinates can be used to combine it with nearby other k-space profiles in the manifold. For reconstructing images from the combined profiles, different weights are assigned to the nearby profiles according to their Euclidean distances (in the manifold) to the current profile. Finally, an image can be reconstructed from the combined and weighted radial profiles using the non-uniform fast Fourier transform (NUFFT) at each of the acquired profile positions. The details of each of these steps are described below.

A. K-space Data Acquisition and Grouping

Data acquisition is performed using a RGA trajectory (Fig. 1 (a)) under free breathing. Compared to Cartesian acquisition, the RGA trajectory is less sensitive to motion [23], which has benefits for dynamic imaging. When acquiring profiles according to the RGA trajectory, the angle between each two consecutive profiles is 111.246°. This enables a uniform coverage of k-space with high temporal incoherence for any arbitrary number of consecutive profiles.

The k-space data are denoted by $X = [x_1, \ldots, x_Z]$, where the columns $x_i$ are k-space profiles that each have $S$ samples (k-space points in the readout direction). All radial profiles are evenly assigned to $G$ overlapping groups according to their profile angle, where each group contains $P$ profiles. The remaining profiles that were acquired after the $K^{th}$ ($K=G\times P/2$) are not used. As illustrated in Fig. 1 (b) left, the dashed black lines define the angular boundaries of the groups. Each profile is a member of two adjacent overlapping groups. We only colour code profiles from three (out of $G$) indicative groups that are correspondingly shown in the manifold in Fig. 1(b) right. Profiles other than those three groups are colour-coded as dark grey. We denote this grouped high-dimensional data as $X_g$, where $g$ ($1 \leq g \leq G$) is the index of the group. Because of the RGA trajectory, the acquisition times of the radial profiles within each group are evenly distributed across the entire image acquisition period. Therefore, all of the groups share data from common respiratory cycles, and can be embedded into a common manifold with a reduced dimensionality of $d$. The use of overlapping groups results in a greater number of profiles for each group and therefore more robust alignment in the MA stage. As the central low frequency region is more important in motion estimation, we calculate a Gaussian weighting (with standard deviation $\sigma_i$) vector for each profile, centred on the central point of the profile, denoted by $v$ in equation (1).

$$v_i = e^{-\frac{(x_i - s/2)^2}{2\sigma_i^2}}$$

(1)

where $s$ is the index for the k-space sample point ($s=1, \ldots, S$ rows of $x$). We assume that the profiles within each group are comparable to each other, since they represent frequency content at approximately the same orientation (but potentially different motion states). However, because the orientations of the profiles are not exactly the same, this assumption may not be valid in high frequency regions of k-space. These Gaussian weights allow lower frequency k-space regions to contribute more than the higher frequency regions in the intra-group comparison. Note that, for a smaller number of groups (i.e. a larger number of profiles per group, $P$), the central overlapping region of the profiles within the same group is smaller, therefore a smaller $\sigma_i$ is chosen. Instead of using the acquired complex data, only the magnitudes of the k-space samples weighted by $v$ are used as the input to the subsequent MA process.

B. Manifold Alignment

The k-space dataset now consists of $G$ groups, where each group contains $P$ profiles and each profile has $S$ samples. The MA method is used to simultaneously reduce the dimensionality ($\mathbb{R}^S$) of each profile and align all groups of profiles into a common manifold, in which profiles acquired at similar respiratory positions are close together (Fig 1 (b)). The dimensionality reduction and alignment of the groups is performed using our recently proposed MA scheme [14] with some adaptations for the application in this paper. We briefly review this technique here. The MA scheme estimates the low dimensional embeddings $Y_g$ of the original data $X_g$ by minimising a joint cost function:

$$\Theta_{total}(Y_1, \ldots, Y_G) = \sum_{g=1}^G \sum_{p} \sum_{q \in \eta(p)} W_{pq}^{(g)} \left| y_p^{(g)} - y_q^{(g)} \right|^2 + \frac{\mu}{2} \sum_{g=1}^G \sum_{m \neq n} \sum_{i,j} U_{ij}^{(nm)} \left| y_i^{(m)} - y_j^{(n)} \right|^2$$

(2)

The first term is the locally linear embedding (LLE) cost function [24], which represents the intra-group embedding errors. LLE forms the low-dimensional manifold by preserving locally linear relations (encoded by the weight matrix $W^{(g)}_{pq}$) derived from the original high-dimensional data $X_g$ for each group. $y_p^{(g)}$ represents the manifold coordinates of profile $p$ in group $g$. When comparing two profiles for the $W^{(g)}_{pq}$ calculation, the weighted $l^2$ norm (weights are calculated as in equation (1)) is used for calculating the Euclidean distance between the two profiles. Subsequently, the relations at the $p^{th}$ high dimensional data vector are represented by a weighted ($W^{(g)}_{pq}$) linear combination of its $K^{LLE}$ nearest neighbours with index $q (q \in \eta(p))$.

The second term in equation (2) represents the inter-group cost function. $y_i^{(m)}$ represents the manifold coordinates of profile $i$ in group $n$. $U_{ij}^{(nm)}$ is an inter-group similarity kernel which will be discussed later. $\mu$ is a weighting parameter that balances the intra-group and inter-group terms.

The total cost function $\Theta_{total}$ can be rewritten in matrix form as
\[ \varphi_{\text{total}} = \text{Tr}(VHV^T) \quad (3) \]

where \( V = [Y_1, \ldots, Y_c] \) is a \( d \times (G \cdot P) \) matrix containing the coordinates of the embeddings. \( \text{Tr}(\cdot) \) is the trace operator. \( H \) is a composition matrix that combines the intra- and inter-term parameters, denoted as

\[
H = \begin{bmatrix}
M^{(t)} + \mu \sum_g D^{(g)} & -\mu U^{(H)} & \ldots & -\mu U^{(G)} \\
-\mu U^{(H)} & M^{(t)} + \mu \sum_g D^{(g)} & \ldots & -\mu U^{(G)} \\
\vdots & \vdots & \ddots & \vdots \\
-\mu U^{(G)} & -\mu U^{(H)} & \ldots & M^{(t)} + \mu \sum_g D^{(g)}
\end{bmatrix}
\]

The diagonal degree matrices \( D^{(nm)} \) are given by \( D^{(nm)}_{ii} = \sum_j U^{(nm)}_{ij} \). The matrices \( M^{(g)} \) are calculated as \( M^{(g)} = (I - W^{(g)})^T (I - W^{(g)}) \), where \( W^{(g)} \) is a matrix that contains the weights \( W^{(g)}_{pq} \) for the LLE term. Under the scaling constraint \( \sum_g U^{(g)} = 1 \), the estimated embeddings \( V \) are given by the second smallest to \( d + 1 \) smallest eigenvectors of \( H \).

In our application, the groups represent k-space profiles acquired at different angles, which are not directly comparable since they represent frequency content for different orientations. A key characteristic of the MA technique in [14], which makes it suitable in our scenario, is that it performs MA without any inter-group comparisons of the original high-dimensional data. This is achieved by forming a graph in which each profile is a node. Each profile is represented by a feature descriptor based on the steady states of random walks in the graph. This feature descriptor, denoted as \( f \), encodes the locations of the nodes (i.e., profiles) within the graph. It enables a robust inter-group profile similarity measurement to be performed based on these feature descriptors via a graph matching method. Furthermore, the shared profiles of two adjacent groups (caused by the overlapping groups, see Fig. 1) enable more robust alignments of the manifolds to be made. Please refer to [14] for detailed descriptions of \( f \). In this paper we extend [14] to introduce an additional measurement term which is based on the temporal positions of the two compared profiles. This allows temporally closer profiles to be embedded closer in the manifold, which results in a more reliable alignment.

The new proposed similarity measurement kernel is defined as,

\[
U^{(nm)}_{ij} = 1 - \left( 1 - e^{-\frac{-\|f^{(n)}_i - f^{(m)}_j\|^2}{2\sigma_2^2}} \right) \left( 1 - e^{-\frac{-\|f^{(n)}_i - f^{(m)}_j\|^2}{2\sigma_2^2}} \right) \quad (4)
\]

In equation (4), the first Gaussian weighted term (with \( \sigma_2 \)) is the similarity measurement proposed in [14], and the second Gaussian weighted term (with \( \sigma_2 \)) is the new temporal weighting measurement. The variable \( t \) represents the acquisition time of the corresponding profile in ms. As an overlapping group structure is used here, every profile is contained in two groups. For a profile that belongs to both groups \( n \) and \( m \) (i.e. when \( \bar{t}^{(n)}_i = \bar{t}^{(m)}_i \)), the value \( U^{(nm)}_{ij} \) is one, which constrains the same profile (or temporally close profiles) from the two groups to be aligned closer. As in [14], the Hungarian algorithm is used to establish one-to-one sparse correspondences between the groups of profiles by maximising the global similarity cost. The resulting one-to-one similarity measurement \( U^{(nm)}_{ij} \) from the Hungarian algorithm is used in equation (2). The final manifold embeddings \( V \) are solved using equation (3).

### C. Image Reconstruction

By embedding all profiles in a common manifold, those that represent similar respiratory positions should be close together. In contrast to conventional gating methods our method does not group the profiles into a limited number of motion states. Our method allows images to be retrospectively reconstructed at as many respiratory positions as the number of k-space profiles (2D) or the number of stacks of k-space profiles (3D) acquired. For each of the acquired k-space profiles, image reconstruction is performed by grouping a number of profiles that have the closest Euclidean distances to the current profile. Duplicated profiles from two groups are only used once. To ensure the selected profiles have an approximately even angular distribution in k-space, the \( L \) highest weighted profiles are selected from 100 evenly divided groups based on the profile radial angles, resulting in \( L \times 100 \) profiles being used for each of the image reconstructions. Note that the 100 groups is a fixed number which is independent of the number of groups \( G \). The NUFFT method [25] is used to reconstruct the final image from the selected radial k-space profiles. In contrast to the conventional NUFFT reconstruction, we use the Euclidean distances between profiles in the manifold to compute weights that determine the contributions of profiles in the non-uniform gridding process. The k-space value \( F(u,v) \) at each grid point \( u \) is calculated as,

\[
F(u) = \sum_{v} e^{\frac{R(u,v)}{2\sigma_2}} F(v) \quad (5)
\]

where \( F(v) \) is the acquired k-space data value at point \( v \) from the candidate radial profiles that contribute to the data re-sampling at \( u \). \( R(u,v) \) are the weighting values that are calculated from the Euclidean distances (in the manifold) of the two profiles \( (y^{(u)} \) and \( y^{(v)} \) that \( u \) and \( v \) belong to

\[
R(u,v) = c(v) \times e^{-\frac{-\|y^{(u)} - y^{(v)}\|^2}{2\sigma_2^2}} \quad (6)
\]

In equation (6), \( c(v) \) is the density compensation weight for the acquired k-space point \( v \), which is calculated in the standard NUFFT process for radial trajectory. \( \sigma_2 \) is set as half of the standard deviation of the embedded manifold coordinates. In [15], the authors used a similar manifold regression method for reconstruction but for k-space lines in a Cartesian trajectory. Using this weighted profile reconstruction scheme, for different profiles, different images are reconstructed even if exactly the same sets of profiles are selected, as the weights of the profiles in each reconstruction would be different.

For 3D image reconstruction of data acquired by the golden radial stack-of-stars method, the Fourier transform is firstly applied along the slice direction. This converts the z-direction encoding from the frequency domain to the image domain, where slice-by-slice reconstruction can be performed. Parameter settings for the free parameters of our method are discussed in section III-B.

### III. MATERIALS

The proposed method was evaluated on both synthetic and in vivo datasets. The 2D and 3D synthetic datasets were used to establish the ground truth to quantitatively evaluate the image reconstruction accuracy of the proposed method. We also
demonstrated the practical feasibility of our technique using in vivo datasets in both 2D and 3D acquisitions, and the results were compared with an adapted version of the well-established CKG method.

A. 2D and 3D Synthetic Dataset Generation

To mimic a realistic data acquisition process, we generated high spatial and temporal resolution 3D sequences, based on image registration of a respiratory gated high spatial resolution (RGHR) 3D MRI volume to a dynamic 3D low spatial resolution (DLR) MRI sequence. The DLR sequence has temporal resolution of ~260 ms and a limited number of motion states (35 in our data). Random volume selection and interpolation were used to generate a realistic ground truth sequence with high temporal resolution (~3 ms). From this, multi-coil k-space data were simulated with the same temporal resolution per k-space profile. An overview of the process is illustrated in Fig. 2, and consists of five steps.

(a) The RGHR volume was transformed to align with the corresponding end-exhale DLR volume using B-spline deformable image registration [26] (Fig. 2 (a)).

(b) The end-exhale DLR volume was registered with all other DLR volumes (Fig. 2 (b)). Therefore, the transformations were established that could deform the aligned RGHR volume to the respiratory positions of each of the DLR volumes.

(c) In order to generate long and realistic synthetic dynamic sequences, the DLR volumes were grouped into four different respiratory groups (end-inhale, end-exhale, mid-inhale, and mid-exhale) according to their diaphragm positions in the central slice. To construct a single random respiratory cycle, we randomly selected (based on a uniform distribution) one DLR volume from each of the four groups. Using their corresponding B-spline transformations, extra DLR volumes were interpolated between these four volumes to produce a high temporal resolution breathing cycle. For the experiments in this paper, different numbers of such interpolated volumes were used to simulate slow and fast breathing cycles. For all cycles, the final synthetic sequences had a temporal resolution of ~3 ms.

(d) The aligned RGHR volume was transformed to each of the respiratory states in each cycle based on the corresponding registered and interpolated B-spline transformations.

(e) Multi-coil (8 coils) images were generated for each volume using coil sensitivities simulated with an analytic integration of Biot-Savart equations. To mimic the 3D golden angle stack-of-stars acquisition [27], only one k-space profile was extracted from a single slice of each volume. K-space profile simulation was performed along the slice direction first before moving to the next golden angle position. For the 2D synthetic dataset, the same procedure was applied with the number of slices set to 1.

The 3D RGHR volume was acquired with respiratory gating at the end-exhale position from a volunteer, with TR = 4.4 ms, TE = 2.2 ms, flip angle = 90°, acquired voxel size 2.19x2.19x2.74 mm³, acquired matrix size 160x160x120, reconstructed voxel size 1.37x1.37x1.37 mm³ and reconstructed matrix size of 256x256x240. The acquisition window was approximately 100 ms, and the scan time was approximately 5 mins. The 3D DLR sequence of the same volunteer contained 35 dynamics under free-breathing using cardiac gating at late diastole to minimise cardiac motion. 3D TFEPI was employed to acquire each volume with TR = 10 ms, TE = 4.9 ms, flip angle 20°, acquired voxel size 2.7x3.6x8.0 mm³, acquired matrix size 128x77x20, reconstructed voxel size 2.22x2.22x4.0 mm³, reconstructed matrix size 144x144x40, TFE factor 26, EPI factor 13, TFE acquisition time 267.9 ms. All the RGHR and DLR volumes were re-sampled into volumes with a voxel size of 1x1x1 mm³ using trilinear interpolation. In this paper, we focus on respiratory motion in the liver-lung region, and to mimic the in vivo acquisition, a region-of-interest (red box in Fig. 2) with voxel size of 1x1x8 mm³ (matrix size 200x200x10) was extracted from the original image.

To make it comparable with the in vivo acquisitions, based on the above process, we generated 9000 multi-coil profile data
for each of the 2D datasets, and 5000 multi-coil stacks-of-profiles with 10 sagittal slices per volume for the 3D synthetic datasets. In total, we generated 6 such randomised sequences for each of the 2D and 3D scenarios. The datasets contained different numbers of breathing cycles ranging from 8 to 20 breathing cycles per minute. Note that, although a number of realistic motion variations are modelled by this process, some artificial motion states may be introduced due to the limitations of the registration algorithm and the interpolation process.

B. 2D and 3D In vivo Dataset Acquisition

2D RGA data of the liver-lung region was acquired on a Philips 1.5T scanner using a 28 channel-coil on five healthy volunteers. Acquisition was performed under free-breathing for approximately 30s, resulting in approximately 9000 k-space profiles for each volunteer. A sagittal balanced SSFP acquisition was performed with 2 mm × 2 mm × 8 mm resolution, FOV=320 mm × 320 mm, flip angle 70°, TR=3.08 ms, TE=1.54 ms. The reconstructed matrix size was 160×160 with a pixel size of 2 mm × 2 mm.

For 3D in vivo datasets, the 3D stack-of-stars with RGA trajectory was employed for the data acquisition in the liver-lung region of five volunteers. Data was acquired on a Philips 1.5T scanner using a 28 channel-coil. All profiles corresponding to one radial angle were acquired sequentially in the slice direction (kz) before moving to the next angle. A sagittal balanced SSFP was performed with TR = 3.8 ms, TE = 1.9 ms, FOV=260 mm × 260 mm × 64 mm, flip angle = 70°, resolution 2.0 mm × 2.0 mm × 8.0 mm, reconstructed matrix size of 176x176x10. There were a total of 5600 stacks of profiles acquired under free-breathing in approximately 5 minutes for each volunteer.

IV. EXPERIMENTS AND RESULTS

A. Comparative Technique

Due to the lack of ground truth for the in vivo datasets, and the fact that the conventional CKG method typically reconstructs images at a limited number of gating windows, we implemented the CKG method with adaptive gating windows to achieve higher temporal resolution reconstruction for comparison. For the 2D dataset, the central k-space magnitudes of all acquired radial profiles were extracted and filtered (1D Gaussian filter with variances of 100 for 2D dataset and 10 for 3D dataset) to estimate a 1D respiratory signal. To reconstruct a gated image at a specific profile location, a gating window (1/20 of the maximum respiratory amplitude) was set around the current profile’s k-space magnitude. Any profiles (\(K_{\text{gating}}\)) with a k-space magnitude within the gating window were assumed to represent similar respiratory positions, and the \(K_{\text{gating}}\) temporally closest profiles were selected for image reconstruction. If \(K_{\text{gating}} < K_{\text{gating}}\), the gating window was iteratively increased by 1/200 of the maximum amplitude until \(K_{\text{gating}} \geq K_{\text{gating}}\). A similar CKG method was applied to the 3D sequence, in which the central k-space magnitudes of the central slice were used to form the 1D respiratory signal. The number of profiles used for image reconstruction was the same as for our MA-based method in order to allow a fair comparison.

B. Evaluation Criteria

Correlation of virtual navigator (CVN): Based on the synthetic dataset, a virtual navigator (VN) [28] was used to measure the head-foot diaphragm translations of the reconstructed and the ground truth sequences. For the 2D dataset, the position of the liver-lung boundary of the sagittal view in the central column of the image was used. The central slice of the 3D dataset was used for the VN measurement by the same 2D method. We calculated the Pearson’s correlation coefficient (PCC) between the VN values measured from the ground truth and the reconstructed sequences as one of the evaluation criteria.

Image intensity correlation (ICC): Normalised cross correlation (NCC) was used to compare the image intensities of the ground truth and the reconstructed sequences for the synthetic datasets. The NCC value was calculated for each corresponding pair of ground truth and reconstructed images and an overall mean and standard deviation of NCC is reported.

Peak Signal to Noise Ratio (PSNR): The PSNR of the reconstructed synthetic dataset was calculated using the corresponding ground truth sequence as the reference, as below:

\[
PSNR = 10 \log_{10}(\frac{\text{peakI}^2}{\text{MSE}})
\]

Where peakI is maximum value of the image’s intensity range. MSE is the mean square error between the reconstructed image and the ground truth image.

Local Sharpness (LS): A sharpness measurement was used as a quantitative measure of image quality for both the synthetic and the in vivo datasets. Five lines were selected at the liver-lung boundary (3 lines), main vessel in the liver (1 line) and rib region (1 line). The sharpness at each line was measured as the maximum of the image gradient magnitude divided by the maximum image intensity of the line [29]. The average sharpness value of the five lines was used as an evaluation criterion for each image. The sharpness values are in the range [0, 1], and sharper structures have higher sharpness values.

Gradient Entropy (GE): For both the synthetic and in vivo datasets, the entropy was calculated based on the gradient magnitude of the image [30]. Lower GE values indicate fewer image reconstruction artefacts.

C. Parameter Tuning

As described in section III-A, we generated 6 synthetic datasets for each of the 2D and 3D cases. For both 2D and 3D, validation was performed using a two-fold cross validation on the 6 datasets, i.e. the free parameters were tuned using one fold and applied to the other fold. The evaluation criterion for parameter tuning was the sum of IIC and CVN.

The performance of our MA-based technique is mainly dependent on the number of profiles per group \(P\) and the number of groups \(G\), where the product of \(P\) and \(G\) is the total number of profiles \(K\). A larger \(P\) enables each group to contain more sampled profiles, which makes manifold embeddings more robust. On the other hand, a larger \(G\) means a smaller angular difference between the sampled profiles within each group, which enables a larger k-space region to be used for a richer comparison of profile data. Hence, a balance between \(P\) and \(G\) needs to be achieved. In order to capture the inter- and intra-cycle variations, a sufficient number of samples per
breathing cycle per group (A) is essential. The number of breathing cycles (B) is automatically determined from the central k-space magnitude. G is then calculated as $[K/P \times 2]$. The weight $\mu$ that balances the intra-group and inter-group terms in equation (1) is determined by the cross validation as described below.

Performance was found not to be sensitive to the remaining parameters, which we determined through a parameter sweep on one 2D and one 3D synthetic dataset. The following settings were consistently applied to both 2D and 3D cases. $K^{\text{LLE}}$ is the number of neighbours used to estimate the local relationships for the LLE, which was set as $[P/10]$ with the minimum of 15. $\sigma_1$ is the Gaussian-weighting for the radial profiles, which was automatically determined by the angular range per group, calculated as $[G/2r]$ with minimum of 3 for cases of very small G. Furthermore, $\sigma_2$ (Gaussian-weighting for similarity kernel) and $\sigma_3$ (temporal constraint in the manifold) were set to 0.1 and 150 ms respectively. The dimensionality of the manifold was set to 3, as suggested in [2]. Finally, L (number of selected profiles from 100 evenly distributed groups for image reconstruction) was $[1 \times \pi/100]$, with the reconstructed image size of $I \times L$.

For the two-fold cross validation, we varied the parameters A and $\mu$ based on three datasets, and applied the optimum A and $\mu$ that achieved the best performance to the remaining three datasets. The process was then switched. The evaluation range of $\mu$ was $[10^{-5}, 10^{-3}, 10^{-5}, 10^{-1}, 10^{0}]$ for both the 2D and 3D cases. The evaluation ranges of A for 2D and 3D tests were $[10, 20, \ldots, 150]$ and $[5, 10, 15, \ldots, 30]$ respectively. Parameter A for 3D datasets had a smaller range because fewer samples were acquired per breath cycle due to the data acquisition along the slice direction.

Table 1. Optimum parameter settings for 2D and 3D synthetic datasets.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Values</th>
<th>Method of tuning</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>80 (2D fold 1) 100 (2D fold 2) 20 (3D fold 1) 25 (3D fold 2)</td>
<td>2-fold cross validation</td>
</tr>
<tr>
<td>$\mu$</td>
<td>$10^{-3}$ (2D both folds) $10^{-3}$ (3D both folds)</td>
<td>2-fold cross validation</td>
</tr>
<tr>
<td>$P$</td>
<td>$A \times B$</td>
<td>Automatically determined</td>
</tr>
<tr>
<td>$G$</td>
<td>$[K/P \times 2]$</td>
<td>Automatically determined</td>
</tr>
<tr>
<td>$K^{\text{LLE}}$</td>
<td>$[P/20]$ with min=15</td>
<td>Parameter sweep</td>
</tr>
<tr>
<td>$\sigma_1$</td>
<td>$[G/2r]$</td>
<td>Parameter sweep</td>
</tr>
<tr>
<td>$\sigma_2$</td>
<td>0.1</td>
<td>Parameter sweep</td>
</tr>
<tr>
<td>$\sigma_3$</td>
<td>150 ms</td>
<td>Parameter sweep</td>
</tr>
<tr>
<td>$d$</td>
<td>3</td>
<td>Literature</td>
</tr>
<tr>
<td>$L$</td>
<td>$[1 \times \pi/100]$</td>
<td>Nyquist sampling theory</td>
</tr>
</tbody>
</table>

The best mean IIC measures of the two folds for both 2D and 3D datasets were the same. The best mean CVN measures of fold 1 and fold 2 for the 2D tests were 0.990 and 0.987 respectively. For the 3D tests, the best mean CVN values for fold 1 and fold 2 were 0.980 and 0.983 respectively. None of the differences were statistically significant. The parameter settings for both 2D and 3D datasets are summarised in Table 1, which were consistently applied in the following evaluation section. For the optimized settings, a typical range of $P$ is 300-600 for 2D and 800-1200 for 3D, which normally resulted in using $\sigma_1$ in the range of 3 to 9.

D. Evaluation Results of 2D and 3D Synthetic Datasets

The two-fold cross validation results for the synthetic datasets are reported in this section. Since a multi-coil acquisition was simulated, the data from only one coil that was most sensitive to the respiratory motion was automatically selected for the MA process. The coil that had the highest spectral magnitude, which was derived from the time-series k-space centre values, in the frequency range of 0.5-2.5 Hz was used [25]. The profiles were then embedded into a common manifold, in which each point represents a profile. A sample embedding is shown in Fig 3 (a). The colours used to visualise the embedded points were determined by the normalised magnitude of the central k-space value, and these values were used only for visualisation purposes and not to determine the embeddings. As can be seen, the proposed MA method successfully embedded profiles that represent similar respiratory positions close together. Furthermore, it has successfully captured both intra- and inter-cycle variations in respiratory motion. More investigations are performed to explore this behavior for the in vivo dataset in the next section. Note also that the points visualised in Fig. 3 (a) represent k-space profiles with a range of different angles, as the manifold is a common space for embedding all angular groups.

For the 3D synthetic dataset with 5000 stacks of profiles, it was assumed that there was no significant motion for the simulated acquisition for a given radial angle along the slice direction (i.e. for one stack of profiles). The only difference from the 2D method was that the stack of profiles at the same radial angle was concatenated to form a single vector to be used by the MA method. The parameter settings described in section III-C were used. An example manifold embedding is shown in Fig. 3 (b).

The entire 2D and 3D sequences were reconstructed using the method described in section II-C. The IIC, CVN, PSNR, LS and GE measurements were used to evaluate the performance of the CKG and the proposed MA methods. An example of a reconstructed 3D dataset and absolute difference images with the corresponding ground truth is shown in Fig. 3 (c). Examples of the VN measurement of a 2D and a 3D synthetic dataset are shown in Fig 4. The numerical results of all 6 synthetic datasets for both 2D and 3D are summarised in Table 2. Except for the CVN in 2D and GE in 3D, all measurements were found to be statistically significantly better using the proposed MA method compared to the CKG method. This calculation was based on a one-tailed Wilcoxon signed rank test using a 99% confidence level.

Table 2. Evaluation results for 2D and 3D synthetic datasets. The values reported in the table are the mean ± standard deviation of all 6 synthetic sequences.

<table>
<thead>
<tr>
<th>Synthetic</th>
<th>2DCKG</th>
<th>2DMA</th>
<th>3DCKG</th>
<th>3DMA</th>
</tr>
</thead>
<tbody>
<tr>
<td>IIC</td>
<td>0.990±0.002</td>
<td>0.996±0.002*</td>
<td>0.991±0.002</td>
<td>0.996±0.003*</td>
</tr>
<tr>
<td>CVN</td>
<td>0.985±0.024</td>
<td>0.989±0.020</td>
<td>0.975±0.027</td>
<td>0.982±0.026*</td>
</tr>
<tr>
<td>PSNR</td>
<td>33.01±2.90</td>
<td>34.98±2.93*</td>
<td>31.69±3.36</td>
<td>34.65±2.42*</td>
</tr>
<tr>
<td>LS</td>
<td>0.178±0.02</td>
<td>0.194±0.02*</td>
<td>0.178±0.02</td>
<td>0.194±0.01*</td>
</tr>
<tr>
<td>GE</td>
<td>1022.5±7.9</td>
<td>1012.94±2.8*</td>
<td>1027.06±22.5</td>
<td>1019.9±10.0</td>
</tr>
</tbody>
</table>

* Statistically significant (p<0.01) improvement of MA over CKG, using one-tailed Wilcoxon signed rank test.
Fig. 3. The embedded profiles in the manifold using the MA method, based on (a) a 2D synthetic dataset with 9000 profiles and (b) a 3D synthetic dataset with 5000 stacks of profiles. Each k-space profile or stack of profiles is represented as a dot in the manifold. The profiles were colour coded using the normalised central k-space magnitude. (c) Example reconstructed images of a 3D synthetic dataset at slice positions 3, 5 and 7, using the MA method (top row). The absolute difference image between the reconstructed image using MA and the ground truth (bottom row).

Fig. 4. (a) Virtual navigator values of the reconstructed sequence of a 2D synthetic dataset and its corresponding ground truth. Pearson’s correlation coefficient is 0.9866. (b) Virtual navigator values of the reconstructed sequence of a 3D synthetic dataset and its corresponding ground truth. Pearson’s correlation coefficient is 0.9813.

E. Evaluation Results of 2D and 3D In vivo Datasets

The proposed method was also evaluated on 2D and 3D in vivo datasets (section III-A) of five healthy volunteers. The parameter settings were the same as the 1st fold of the synthetic dataset (A=80 for 2D and A=20 for 3D). Also, as with the synthetic data, the profiles from the coil that was most sensitive to the respiratory motion were used for the MA process. Subsequent image reconstruction was performed using data from all coils. The normalised magnitude of the centre of k-space (Fig. 5 (a)) was used to colour code the embedded points. In Fig. 5 (a) the first mode of the manifold obtained using the MA method is also plotted for comparison, which correlates (Pearson correlation coefficient of 0.98) well with the k-space magnitude. The 3D manifold embedding of a sample 2D in vivo dataset is shown in Fig. 5 (b). Similar to the synthetic dataset, the multi-dimensional manifold embedding of the 2D in vivo dataset captures significant amounts of intra- and inter-cycle variation. In Fig. 5 (c), we show the same embedding as Fig. 5 (b) but using a different colour coding which shows inspiration and expiration profiles using blue and red respectively. It is clearly observed that the inhale and exhale profiles are automatically distinguished by the 3D manifold embedding. This represents the well-known hysteresis effect, in which the motion states passed through during inspiration are different to those passed through during expiration [31]. The entire dynamic sequence of the 2D in vivo dataset was reconstructed. Since the ground truth is unknown for the in vivo datasets, the CKG method (section IV-A) was used for comparison purposes. Fig. 6 shows example images that were reconstructed by the CKG method (top row) and our MA method (bottom row) at four different respiratory positions. Less motion and radial streaking artefacts can be observed with the proposed method, especially at small structures.

Table 3. Evaluation results for 2D and 3D in vivo datasets. The values reported in the table are the mean ± standard deviation over all 5 in vivo datasets.

<table>
<thead>
<tr>
<th>In vivo</th>
<th>2DCKG</th>
<th>2DMA</th>
<th>3DCKG</th>
<th>3DMA</th>
</tr>
</thead>
<tbody>
<tr>
<td>LS</td>
<td>0.37±0.03</td>
<td>0.41±0.04*</td>
<td>0.34±0.04</td>
<td>0.38±0.06*</td>
</tr>
<tr>
<td>GE</td>
<td>570.75±19.33</td>
<td>543.44±15.60*</td>
<td>571.31±19.59</td>
<td>544.19±21.32*</td>
</tr>
</tbody>
</table>

* Statistically significant (p<0.01) improvement of MA over CKG, using one-tailed Wilcoxon signed rank test.

A sample reconstruction of a 3D in vivo dataset is shown in Fig. 7, with slices shown at the 3rd, 5th and 7th slice positions and also a volume rendering using maximum intensity projection. By comparing the reconstructed images between the CKG and the MA methods, it is clearly seen that better image quality with fewer artefacts was achieved by the MA method. The manifold embeddings (Fig. 7 leftmost image in bottom row) captured larger variations in the end inspiration states than the end expiration states, which is consistent with the observations from the central k-space magnitude (Fig. 7 leftmost image in top row). Numerical results of the LS and GE measurements of the CKG and the MA methods for both 2D and 3D in vivo datasets were compared against the ground truth using Wilcoxon signed rank test. The proposed method showed statistically significant improvements (*p<0.01) over the CKG method for both 2D and 3D datasets.
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Datasets are reported in Table 3. The numbers are means and standard deviations over the reconstructed sequences of the 5 in vivo datasets. Both the LS and GE measurements show a statistically significant improvement using the MA method.

**V. DISCUSSION AND CONCLUSION**

We have presented a novel technique for retrospective dynamic MRI reconstruction, based on MA. The method enables...
reconstruction of ‘motion-free’ abdominal images throughout entire respiratory cycles. Our method was evaluated on both 2D and 3D synthetic and in vivo datasets. We have shown both visually and quantitatively the improvement of using the MA method against the CKG method. Statistically significant improvements were found in the numerical comparisons, in terms of image intensity, virtual navigator measurement, PSNR, local sharpness and gradient entropy.

Our proposed technique has a number of advantages compared to the current state-of-the-art in MRI self-gating. First, the use of MA permits much richer profile data to be used for establishing respiratory state correspondences for self-gating purposes. This provides an advantage over the CKG method by extracting a multidimensional signal (rather than the 1D signal), which captures more dimensions of the motion. Specifically, our technique uses a Gaussian-weighted version of the entire profile rather than only the central value as in CKG. This is made possible because our manifold embedding process only ever compares k-space data within a group, where all data are comparable and differences will be due to motion only. The use of richer profile data in the embedding process permits better estimation of underlying intra-cycle and inter-cycle variation in respiratory motion, as can be seen from the separation of inspiration and expiration profiles in Fig. 5 (b) and (c). This separation was consistently observed for all in vivo datasets. However, it is worth noting that inspiration and expiration are only separable in the manifold embeddings when large enough numbers of samples per breathing cycle are acquired (e.g. as in the 2D acquisition). It is more difficult to capture such variations in the 3D acquisition (Fig. 3 (b) and Fig. 7), because the greater time taken to acquire a stack of profiles means that fewer such stacks are acquired in each breathing cycle.

Another key contribution of our approach is that we perform 3D MRI MA self-gating directly using the k-space data. To the authors’ knowledge, all manifold-learning-based self-gating techniques to date have worked in 2D only [15, 18, 20]. Those techniques would be non-trivial to extend to 3D. Previous work in [15] and [18] involve a repetitive acquisition of certain k-space lines either in a Cartesian or radial trajectory. Therefore, significant motion is likely to occur during the additional acquisition along the slice direction in 3D. With the RGA acquisition in [20], the intersection region of a number of consecutive spokes was used for the manifold embedding. To extend this technique to 3D would require a multiple 2D slice by slice acquisition, and the motion state correspondences between slices would be unknown. To address this issue, we have recently proposed a different MA framework for multi-slice acquisitions [16].

Our results show that impressive reconstructions can be achieved in both 2D and 3D. The weighted profile image reconstruction scheme efficiently uses profiles from other nearby motion states, which potentially reduces the total number of profiles to be acquired, when compared to conventional CKG. The computational time for MA is proportional to the total number of profiles. For 5000 stacks-of-profiles in 3D, it takes about 60 minutes to run the MA on a 3.6 GHz processor with non-optimised MATLAB code. The majority of the time is occupied by the Hungarian algorithm for establishing inter-group correspondences. The image reconstruction time is similar to a standard NUFFT reconstruction time. We modified the NUFFT code (without CS) in [25] with our weighted profile reconstruction. It takes about 50 seconds to reconstruct a 176x176x10 voxels 3D volume, which is significantly less than the CS method. However, we do not view our MA method as an alternative to CS techniques, but rather a complementary approach.

As for other self-gating methods, our MA method cannot achieve a good reconstruction if an insufficient number of profiles is available at similar respiratory positions. This would happen if, for example, an extreme inhale position was visited relatively infrequently during the imaging period. In future work we plan to address this problem by investigating if the extreme positions could be detected by calculating the variations of the weights that contributed to the reconstruction. Higher variation than a pre-defined threshold could indicate a higher likelihood of poor reconstruction quality, so this information could be used to trigger reconstruction using an alternative but more time consuming technique like CS.

Another area for future work is to extend our method to other sampling trajectories. We believe that our MA-based framework could be applied to Cartesian sampling trajectories. In this case, groupings of k-space profiles would be based on Cartesian rows rather than radial angles. Work is also in progress to extract signals from data that are corrupted by both cardiac and respiratory motions.

In conclusion, we believe that our proposed MA-based self-gating method for dynamic MRI represents an improvement on the current state-of-the-art, and a novel application for MA in MRI imaging. The better quality and considerably higher temporal resolution reconstructed dynamic images will be useful in a range of applications such as motion-correction of PET data in a simultaneous PET-MR scenario.

DATA DOWNLOAD
The data and code for generating the synthetic datasets are available at: http://kclmmag.org/downloads.html.

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REFERENCES


