

High-Resolution Motion-Robust Diffusion-Weighted Imaging with Self-Gated Zero-Shot Self-Supervised Reconstruction

Zhengguo Tan, Patrick A Liebig, Annika Hofmann, Frederik B Laun, Florian Knoll

Abstract— This work introduced a self-gated zero-shot self-supervised learning (ZSSL) reconstruction framework for navigator-free, high-resolution diffusion-weighted imaging using undersampled multi-shot interleaved echo-planar imaging (iEPI) acquisition. ZSSL belongs to the algorithm unrolling technique, with a physics-guided data-consistency term and a learned regularization function. We unrolled the alternating direction method of multipliers (ADMM) with a residual neural network to leverage redundancy across the spatial-diffusion-dimension. First, we compared the proposed self-gated ZSSL to conventional methods including parallel imaging as multiplexed sensitivity-encoding (MUSE) and compressed sensing reconstruction with locally-low rank (LLR) regularization. ZSSL supplied excellent reconstruction results in both 4-shot fully-sampled data and 2-shot undersampled data at 1.0 mm isotropic resolution. Second, we demonstrated the capability of self-gated ZSSL in DWI reconstruction with the b-value of 3000 s/mm². Compared to LLR, ZSSL provided more noise reduction and clearer diffusion contrast. Third, self-gated ZSSL was validated with both retrospectively and prospectively acquired data at 0.7 mm isotropic resolution. This approach outperformed both MUSE and LLR regularized reconstruction in terms of image sharpness and motion robustness. While ZSSL required up to eight hours training time per slice, it generalized well to all other slices, with an inference time of just one minute. In comparison, LLR required about two hours per slice. Overall, self-gated ZSSL enables undersampled multi-shot iEPI acquisition without the need of navigators, providing sub-millimeter DWI at clinically feasible reconstruction times. The code is publicly available at: <https://github.com/ZhengguoTan/DeepDWI>.

Index Terms— Diffusion weighted imaging, Magnetic resonance imaging, Image reconstruction, End-to-end learning in medical imaging, Machine learning

I. INTRODUCTION

HIGH-dimensional magnetic resonance imaging (HD-MRI) has been a flourishing field, focused on the acquisition, reconstruction and analysis of multi-dimensional multi-contrast-weighted MRI data. Examples of HD-MRI include but are not limited to magnetic resonance spectroscopic imaging (MRSI) [1], diffusion-weighted imaging (DWI) [2], [3], and quantitative parameter mapping [4], [5]. Conventional HD-MRI, however, requires long acquisition times, making the data susceptible to subject motion and system imperfections, and imposing high computational burden. DWI, in particular, poses challenges in the pursuit of high spatial, temporal, and angular resolution. DWI is typically acquired using the pulsed gradient spin echo sequence [6] followed by fast echo-planar imaging (EPI) readouts [7]. However, the use of long echo trains in EPI results in geometric distortion artifacts and reduced spatial resolution. Additionally, acquiring multiple diffusion directions to enhance angular resolution and to better probe tissue microstructure further extends the scan time.

Advances in parallel imaging [8]–[12] and compressed sensing [13]–[15] have enabled accelerated acquisition for HD-MRI. Notably, the low-rank model [16] has been a powerful tool in dimension reduction. Typically, singular value decomposition (SVD) is used to learn a truncated temporal basis function from a large-scale physics-informed dictionary [17]–[19]. The temporal basis function is then integrated with the MRI forward model, i.e. the sensitivity encoding operator [11], for joint reconstruction of the corresponding spatial basis images. In addition, low-rank regularization can be employed in the joint reconstruction [20].

Beyond the low-rank technique, advanced neural networks, e.g. autoencoder [21], have been explored for HD-MRI reconstruction and proven to supply more accurate representations of high-dimensional data than SVD. Lam et al. [22] and Mani et al. [23] proposed to first learn a denoising autoencoder (DAE) model from a physics-informed simulated dictionary and then incorporate the learned DAE model as a regularizer in the alternating direction method of multipliers (ADMM) [24] unrolling reconstruction. Pioneered by Gregor and LeCun

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[25], algorithm unrolling enables the use of learned deep *priors* as regularization and offers faster inference compared to iterative reconstruction methods that rely on hand-crafted regularization functions [26]. Algorithm unrolling has been applied to accelerated MRI reconstruction in various scenarios: including but not limited to supervised learning with fully sampled reference images [27], [28], and self-supervised learning with only undersampled data available for training [29], [30]. Notably, acquiring fully sampled DWI for training a regularization function is quite challenging. First, fully-sampled DWI requires longer echo times in EPI, which both elongates the scan times and increases off-resonance-induced geometric distortions. Second, the variety of diffusion acquisition modes necessitates a larger dataset compared to two-dimensional imaging scenarios [31]. As a result, self-supervised learning is better suited for DWI reconstruction.

Deep neural networks are capable of learning not only regularization functions, but also MR-physics forward operators. Liu et al. [32] proposed the reference-free T_1 parameter maps extraction (RELAX) self-supervised deep learning reconstruction, which learns the mapping from T_1 parameter maps to undersampled multi-coil multi-contrast k -space data. Arefeen et al. [33] proposed to replace the conventional SVD-based linear subspace modeling [17] by the latent decoder model within DAE for improved T_2 -weighted image reconstruction. The ability of DAE to learn DWI models is somewhat uncertain. DAE is composed of sequential fully connected layers with nonlinear activation functions, which may struggle with complex functions like those required for DWI signals. For instance, the standard diffusion tensor model [34] consists of six tensor elements, and generates DWI signals based on the multiplication of exponential functions, a process that may be too intricate for simpler architectures to capture effectively.

Contributions:

- We unrolled ADMM to perform zero-shot self-supervised learning (ZSSL) and incorporated self-gated shot-to-shot phase variation estimation into ZSSL for deep diffusion-weighted imaging reconstruction.
- We achieved navigator-free high-resolution DWI with 21 diffusion-encoding directions at 0.7 mm isotropic resolution, and a scan time of under 10 minutes.

II. RELATED WORK

A. Multi-Band Multi-Shot DWI Acquisition & Modeling

Our previous work [35] demonstrated the joint k - q -slice forward operator for multi-band multi-shot navigator-based interleaved EPI (NAViEPI) DWI acquisition. This operator can be understood as an extended sensitivity encoding (SENSE) operator [11], which maps the multi-slice multi-diffusion-weighted images (\tilde{x}) to their corresponding k -space,

$$\mathcal{A}(\tilde{x}) = \mathbf{P}\Sigma\Theta\mathbf{F}\mathbf{S}\Phi\tilde{x} \quad (1)$$

Here, the images \tilde{x} are point-wise multiplied with the pre-computed shot-to-shot phase variation maps (Φ) and coil sensitivity maps (\mathbf{S}). The output images are then converted to k -space via two-dimensional fast Fourier transform (\mathbf{F}), point-wise multiplied with the multi-band phases (Θ), summed

along the slice dimension (Σ), and then multiplied by the k -space undersampling mask (\mathbf{P}).

In Equation (1), one challenge is to accurately estimate the shot-to-shot phase variation. Multiplexed sensitivity-encoding (MUSE) type reconstruction techniques [36]–[39] realized the self-gating strategy, where the k -space data of each shot was used to reconstruct its corresponding shot image followed by a phase smoothing approach. Self-gated shot phase estimation does not require the acquisition of phase navigator data. However, it requires marginal undersampling factors per shot and fully-sampled DWI acquisition assembling all shots. Alternatively, undersampled DWI acquisition can be enabled via the acquisition of navigators for shot phase estimation [35]. This approach allows for mesoscale-resolution DWI at 7 T, but still needs long scan time. As listed in Table I, the total acquisition of Protocol #3 at 0.7 mm isotropic resolution takes 16 : 27 minutes with phase navigators. This scan time can be reduced to approximately 10 minutes by removing the phase navigators (Protocol #4 in Table I).

With the operator \mathcal{A} , the joint reconstruction is expressed as,

$$\underset{\tilde{x}}{\operatorname{argmin}} \|\mathbf{y} - \mathcal{A}(\tilde{x})\|_2^2 + \lambda \mathcal{R}(\tilde{x}) \quad (2)$$

where \mathbf{y} is the measured k -space data. The first term in Equation (2) presents data consistency, and the second term presents the regularization function $\mathcal{R}(\tilde{x})$ with the regularization strength λ . When using the Tikhonov regularization, i.e. $\mathcal{R}(\tilde{x}) = \|\tilde{x}\|_2^2$, Equation (2) can be solved via the conjugate gradient (CG) method. For nonlinear regularization functions, such as the locally-low rank (LLR) regularization [35] or neural networks with nonlinear activation functions, ADMM was employed in this work to solve for Equation (2).

B. Algorithm Unrolling for Deep Image Reconstruction

Algorithm unrolling has been an emerging technique in solving inverse problems with learnable deep neural networks. Algorithm unrolling consists of two ingredients. First, it uses deep neural networks to learn regularization function. Second, it is constrained by the data-consistency term. In other words, the forward pass of the estimate $\mathcal{A}(\tilde{x})$ must be close to the measured data \mathbf{y} . By mapping the operations used in iterative algorithms onto networks, unrolled algorithms can be trained with data, leading to much faster inference than conventional iterative algorithms [26]. Further, recent developments have shown that the operations used in compressed sensing MRI, i.e., sparsifying transformation and soft thresholding, can be learned via neural networks. For instance, Hammernik et al. [27] proposed to unroll the gradient descent algorithm with a learned neural network (e.g. U-net [40]) as the regularization function. Aggarwal et al. [28] proposed the model-based deep learning architecture for inverse problems (MoDL) to unroll the alternating minimization algorithm with a learned residual denoising network [41] as regularization.

C. Self-Supervised Learning for Image Reconstruction

In many MRI applications, such as dynamic imaging and diffusion-weighted imaging acquiring fully-sampled data for

supervised learning can be challenging. To tackle this issue, Yaman et al. [29] proposed self-supervised learning via data undersampling (SSDU), which learns the regularization function in Equation (2) by splitting available undersampled data into two disjoint sets, one of which is used in the data consistency term and another used for the computation in the training loss function. The training of SSDU requires large undersampled data sets. To close the domain gap between training and test data, Yaman et al. [30] proposed scan-specific zero-shot self-supervised learning (ZSSSL), which splits a single data set into three disjoint sets for (a) the data consistency term, (b) the loss calculation during training, and (c) validation, respectively. Recently, ZSSSL has been adopted for multi-contrast image reconstruction [42].

III. METHODS

A. Data Acquisition

Table I lists three acquisition protocols implemented on a clinical 7 T MR system (MAGNETOM Terra, Siemens Healthineers, Erlangen, Germany) with a 32-channel head coil (Nova Medical, Wilmington, MA, USA) and the XR-gradient system (maximum gradient strength 80 mT/m and a peak slew rate 200 T/m/s). Protocols #1 and #2 realized mesoscale DWI with 0.7 mm isotropic resolution. Two-fold acceleration was employed in both in-plane and slice directions. Every DWI was acquired by three shots in an interleaved manner, which resulted in 6×2 -fold acceleration per shot. Noteworthy, the total scan time can be reduced to about 10 minutes (Protocol #2) when switching off navigator acquisition. Protocol #3 realized three-shell diffusion acquisition with b -values spanning 1000, 2000, and 3000 s/mm² at 1.0 mm isotropic resolution. Four young healthy volunteers with written informed consent approved by the local ethics committee participated in this study.

B. Image Reconstruction via ADMM Unrolling and Zero-Shot Self-Supervised Learning

Instead of the two-step alternating minimization unrolling scheme as used in MoDL [28], we employed the ADMM unrolling to solve the self-supervised learning reconstruction in Equation (2). The update rule of ADMM unrolling reads

$$\begin{cases} \tilde{\mathbf{x}}^{(k+1)} = \underset{\mathbf{x}}{\operatorname{argmin}} \|\mathbf{y} - \mathcal{A}(\mathbf{x})\|_2^2 + \rho/2 \|\mathbf{x} - \mathbf{v}^{(k)} + \mathbf{u}^{(k)}\|_2^2 \\ \mathbf{v}^{(k+1)} = (\lambda/\rho) \cdot \mathcal{D}_\omega(\tilde{\mathbf{x}}^{(k+1)} + \mathbf{u}^{(k)}) \\ \mathbf{u}^{(k+1)} = \mathbf{u}^{(k)} + \tilde{\mathbf{x}}^{(k+1)} - \mathbf{v}^{(k+1)} \end{cases} \quad (3)$$

ADMM updates the variables $\tilde{\mathbf{x}}$, \mathbf{v} , and \mathbf{u} in an alternating scheme. It splits the unrolled reconstruction into three steps, as shown in Equation (3) and in the pseudo code of Algorithm 1. First, the updating step for $\tilde{\mathbf{x}}$ is solved by conjugate gradient. Second, the variable \mathbf{v} is then updated via the forward pass of the neural network \mathcal{D}_ω with the input as the sum of current estimates of $\tilde{\mathbf{x}}$ and \mathbf{u} . Third, the variable \mathbf{u} is updated by adding its current estimate to the difference between $\tilde{\mathbf{x}}$ and \mathbf{v} .

As shown in Figure 1, the data sampling mask \mathbf{P} in ZSSSL [30] is split into three disjoint sets, the training mask \mathbf{T} for the

Algorithm 1 ADMM Unrolling for ZSSSL

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1: Initialization:
2:   split sampling mask  $\mathbf{P}$  into 12 repetitions, each of which
   consists of three disjoint sets  $\mathbf{T}$ ,  $\mathbf{L}$ , and  $\mathbf{V}$ 
3:    $p \leftarrow 0$  and  $N_{\text{epoch}} \leftarrow 100$ 
4:    $\mathcal{D}_\omega$  set as ResNet
5:    $\rho \leftarrow 0.05$  and  $\lambda \leftarrow 0.05$ 
6:    $\text{Loss}_{\text{valid}} \leftarrow \inf$  and  $\text{trace} \leftarrow 0$ 
7: function ADMM(mask)
8:    $\mathcal{A}_{\text{mask}} \leftarrow$  set the mask in the forward operator  $\mathcal{A}$ 
9:    $\tilde{\mathbf{x}}^{(0)} \leftarrow \mathcal{A}_{\text{mask}}^H(\mathbf{y})$ 
10:   $\mathbf{v}^{(0)} \leftarrow \tilde{\mathbf{x}}^{(0)}$  and  $\mathbf{u}^{(0)} \leftarrow \mathbf{0}$ 
11:   $k \leftarrow 0$  and  $N_{\text{unroll}} \leftarrow 8$ 
12:  while  $k < N_{\text{unroll}}$  do
13:     $\tilde{\mathbf{x}}^{(k+1)} \leftarrow$  conjugate gradient with 6 iterations
14:     $\mathbf{v}^{(k+1)} \leftarrow (\lambda/\rho) \cdot \mathcal{D}_\omega(\tilde{\mathbf{x}}^{(k+1)} + \mathbf{u}^{(k)})$ 
15:     $\mathbf{u}^{(k+1)} \leftarrow \mathbf{u}^{(k)} + \tilde{\mathbf{x}}^{(k+1)} - \mathbf{v}^{(k+1)}$ 
16:     $k \leftarrow k + 1$ 
17:  end while
18:  return  $\tilde{\mathbf{x}}^{(k+1)}$ 
19: end function
20: Training:
21: while  $p < N_{\text{epoch}}$  or  $\text{trace} \leq 12$  do
22:    $\mathbf{x}_t \leftarrow \text{ADMM}(\mathbf{T})$ 
23:    $\text{Loss}_{\text{train}} \leftarrow \mathcal{L}(\mathbf{Ly}, \mathcal{A}_{\mathbf{L}}(\mathbf{x}_t))$ 
24:   update  $\omega$  via ADAM
25:   Validation:
26:    $\mathbf{x}_t \leftarrow \text{ADMM}(\mathbf{T} \cup \mathbf{L})$ 
27:    $\text{Loss}_{\text{temp}} \leftarrow \mathcal{L}(\mathbf{Vy}, \mathcal{A}_{\mathbf{V}}(\mathbf{x}_t))$ 
28:   if  $\text{Loss}_{\text{temp}} \leq \text{Loss}_{\text{valid}}$  then
29:      $\text{Loss}_{\text{valid}} \leftarrow \text{Loss}_{\text{temp}}$ 
30:      $\text{trace} \leftarrow 0$ 
31:   else
32:      $\text{trace} \leftarrow \text{trace} + 1$ 
33:   end if
34: end while

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data consistency term, the training loss mask \mathbf{L} for the loss function calculation, and the validation loss mask \mathbf{V} . Each set consists of 12 repetitions constructed via random uniform sampling of the data mask \mathbf{P} . In each training epoch, every repetition is looped through in order to update the ResNet parameters ω . Plus, the validation step is performed after every training epoch to update the minimal validation loss. If the validation loss does not reduce for 12 consecutive epochs or if 100 epochs are reached, the training is terminated.

The index k in Equation (3) denotes the unrolling iteration, and \mathcal{D}_ω denotes the residual network (ResNet) [41] parameterized by ω . In this work, 2D convolution was employed to construct the ResNet. In PyTorch, 2D convolution requires four-dimensional tensors as input and output. For instance, a matrix with the size (N, C, H, W) is acceptable for the 'conv2d' function in PyTorch. Here, W and H denote the width and height of the convolution kernel, C denotes the number of channels, and N denotes the batch size. However, the DWIs ($\tilde{\mathbf{x}}$) to be reconstructed has the size $(N_{\text{diff}}, N_Z, N_Y, N_X, 2)$, where 2 stands for the real and imaginary part of the complex-

TABLE I
IEPI WITH OR WITHOUT NAVIGATORS ACQUISITION PROTOCOLS

Protocol	#1 / #2	#3
Diffusion mode		MDDW
Diffusion scheme		monopolar
Diffusion direction	20	114
b -value (s/mm 2)	1000	3-shell
b_0	1	12
FOV (mm 2)		200
Matrix size	286 × 286	200 × 200
In-plane resolution (mm 2)	0.7 × 0.7	1.0 × 1.0
Slice thickness (mm)	0.7	1.0
Slices	176	114
Navigator	Yes / No	No
Shots	3	2
TR (ms)	15000 / 8900	4900
TE (ms)	58/98.3 / 58	70
ESP (ms)	1.17	1.02
Bandwidth (Hz/Pixel)	972	1086
Partial Fourier	5/8	6/8
Acceleration	2 × 2	3 × 3
Acquisition (min)	16 : 27 / 9 : 57	20 : 55

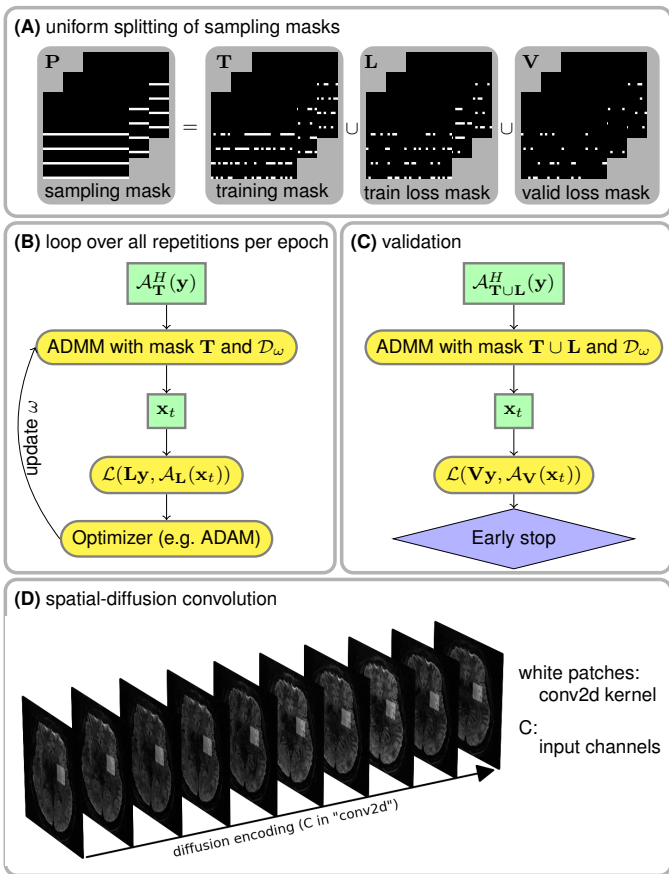


Fig. 1. Illustration of the key components in ZSSSL. (A) The sampling mask P in Equation (1) was uniformly split into three disjoint sets: the training mask T used for the data consistency term during ZSSSL training, the train loss mask L used for the loss function calculation during ZSSSL training, and the validation loss mask V used for the loss function calculation during ZSSSL validation. (B) and (C) show the flowchart for the training and the validation of ZSSSL, respectively. Note that the ResNet parameters ω are updated during training, but remain fixed during the validation step. (D) A stack of DWIs is input into ResNet during ADMM unrolling.

valued DWIs, N_X and N_Y are the width and the height of DWIs, N_Z is the number of slices (same as the multi-band factor), and N_{diff} is the number of diffusion encodings. To train a ResNet based on 2D convolution, the DWIs were reshaped and permuted as $(N_Z, 2 \cdot N_{\text{diff}}, N_Y, N_X)$, as illustrated in Figure 1 (D). In this manner, 2D convolution kernels in combination with ReLU activation functions loop through the varying diffusion-weighted contrast to learn the key features of the high-dimensional data and to reduce noisy and aliasing artifacts in unrolled reconstruction.

C. Comparison of Regularization Techniques

This work compared the reconstruction performance of three different regularization techniques, Tikhonov ℓ^2 regularization (as used in MUSE), LLR regularization, and ZSSSL with a learned regularization. Note that MUSE is a simultaneous multi-slice (SMS) parallel imaging method and poses no regularization along the diffusion dimension, effectively solving each DWI reconstruction independently. In contrast, all other regularized reconstructions fall into the joint reconstruction regime. They jointly reconstruct all DWIs and imposes regularization terms that explore spatial-diffusion redundancy. For example, LLR enforces low rankness of local spatial-diffusion matrices from DWIs, whereas ZSSSL learns a ResNet regularization function based on spatial-diffusion convolution kernels while enforcing data consistency during the unrolled training process.

D. Self-Gated ZSSSL

As discussed in Section II-A, there are two approaches for estimating shot-to-shot phase variation: self-gated and navigator-based. The self-gated approach, as used in MUSE [38], requires fully-sampled DWI acquisition and has typically reported only a small number of shots (up to 4). The previously proposed NAViEPI approach enabled high-resolution DWI with the use of undersampled iEPI and shot-to-shot phase navigator acquisition. While NAViEPI results in shorter scan

time than fully-sampled iEPI, the use of phase navigator still elongates the acquisition, as listed in Table I. Therefore, a key question is whether it would be feasible to discard phase navigator while keeping undersampled iEPI acquisition. In this work, we investigated the feasibility of ZSSSL in self-gated scan for 0.7 mm isotropic resolution DWI.

E. ZSSSL Model Generalization

We evaluated the generalization of the ZSSSL model in two aspects. First, using the 4-shot fully-sampled data acquired by Protocol #1 Table I, we trained ZSSSL with all 4 shots and then tested the trained model with retrospectively undersampled 2-shot data. Second, with the 0.7 mm isotropic resolution DWI data containing 88 multi-band slices acquired from Protocol #3 in Table I, we trained the ZSSSL with only one slice and then performed the inference reconstruction on the remaining slices.

F. Computation

All reconstructions were in this work done on a single A100 SXM4/NVLink GPU with 80 GB memory (NVIDIA, Santa Clara, CA, USA). Computing infrastructure was provided by the Erlangen National High Performance Computing Center.

Note that the data from Protocol #2 in Table I contains a total of 126 diffusion-weighted images, which is too large for the LLR and the ResNet computation in a single A100 GPU. As a result, this data was uniformly split into three consecutive parts, with each part containing 42 diffusion-weighted images.

IV. RESULTS

A. Retrospectively Self-Gated ZSSSL

Figure 2 demonstrates the efficacy of the self-gated ZSSSL reconstruction by comparing to the navigated reconstructions using the NAViEPI data of the first volunteer from Protocol #1 in Table I. The diffusion encoding with accidental inter-shot motion reconstruction results were displayed.

The selected DWIs showed residual aliasing-like and severe motion-blurring artifacts in the navigated reconstructions. The main reason of these artifacts is that the addition of navigator acquisition increased the total scan time, resulting in higher sensitivity to accidental inter-shot motion. Admittedly, navigators are valuable in the case of ultra high resolution using many shots, e.g. 3-fold in-plane undersampling and 5-shot acquisition for the in-plane resolution of 0.5 mm [35]. In this experiment, however, the utilization of 3 shots yielded 6×2 -fold acceleration per shot (refer to Protocol #3). Such an acceleration rate proved achievable in the self-gated approach, as demonstrated in the self-gated reconstruction results. The self-gated ZSSSL reconstruction largely reduced the residual aliasing and motion artifact. Further, self-gated ZSSSL exhibits much clearer tissue delineation in reconstructed DWIs, as indicated by red arrows in the zoomed-in views in Figure 2, whereas self-gated LLR suffers from blurry tissue boundaries and signal dropouts.

Figure 3 shows diffusion-weighted images at the same diffusion encoding as in Figure 2 in the coronal and the sagittal

view, respectively. As mentioned in Section III-E, ZSSSL was trained using only one slice and then inferred on the remaining slices. The ZSSSL model generalized well across slices. The inference of every slice took only about 1 minute, whereas the LLR reconstruction took about 48 minutes per slice. More importantly, the self-gated LLR reconstruction exhibited residual motion-induced stripping artifacts in both coronal and sagittal views [43], whereas the self-gated ZSSSL approach substantially removed these artifacts and supplied high-quality DWI without the need of navigators.

B. Prospectively Self-Gated ZSSSL

Figure 4 compares the reconstruction results using the prospectively acquired iEPI data without navigators of the second volunteer (refer to Protocol #2 in Table I). The snapshot single diffusion-direction DWIs as well as mean DWIs at three orthogonal planes were displayed. The MUSE reconstruction suffered from strong noise at such high spatial resolution.

The LLR regularized reconstruction largely reduced noise, but still exhibited signal void in the axial view, which appeared as striping artifacts in the coronal and sagittal views (refer to the red arrows in Figure 4). These artifacts can be reduced by incorporating adaptive noise estimation in LLR regularization [44]. On the other hand, the LLR regularized reconstruction showed residual noise in the cerebellum region (refer to the blue arrows in Figure 4), which could be caused by the B_1 excitation field inhomogeneity at 7 T.

The above-mentioned artifacts were nearly gone in the ZSSSL reconstruction. The DWI in the axial view showed clearer diffusion contrasts and thus better tissue delineation in the coronal and sagittal views. Plus, the DWI in the sagittal view showed more homogeneous signal distribution.

Figure 5 displays the training and validation loss as well as the learned regularization strength along epochs. It can be seen that 100 epochs were sufficient for the convergence of ADMM unrolling. In addition, the regularization strength converged to the value of about 0.027.

Figure 6 showed the reconstructed DWIs at four diffusion encodings based on the iEPI data acquired from the third volunteer. In this experiment, the volunteer was instructed to keep still during scan. Again, the proposed ZSSSL reconstruction with spatial-diffusion convolution illustrated superior tissue structure delineation to the LLR regularized reconstruction.

C. DWI at b-Value of 3000 s/mm²

Figure 7 compares the LLR and ZSSSL reconstruction results of the 3-shell 1.0 mm isotropic resolution DWI, as acquired by Protocol #3 in Table I. In the single-direction DWI at the b -value of 3000 s/mm², the proposed self-gated ZSSSL showed slight improvement in SNR and structural sharpness compared to LLR. The colored fractional anisotropy maps, as obtained via diffusion tensor model fitting over DWIs in 126 directions, showed similar quality between LLR and ZSSSL.

V. DISCUSSION

This work reported a novel self-gated zero-shot self-supervised learning approach for multi-shot undersampled

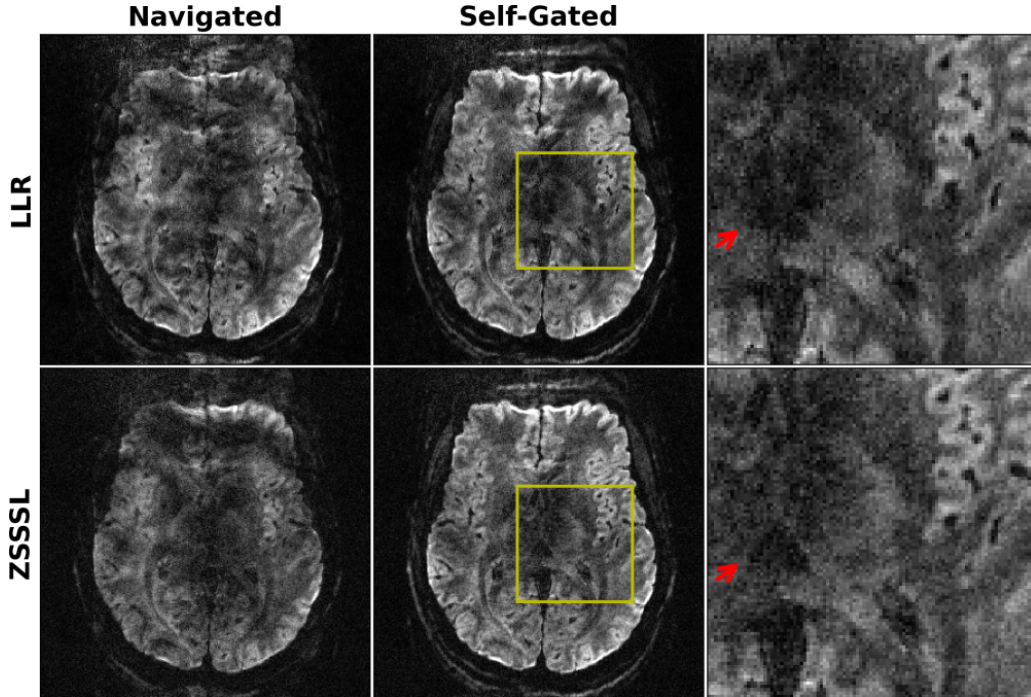


Fig. 2. Comparison of (top) LLR regularized and (bottom) ZSSSL reconstruction on 0.7 mm isotropic resolution DWI acquired by Protocol #1 with shot phase estimated from (left) navigators and (middle) imaging echoes, respectively. Zoomed views of the yellow boxes from the self-gated reconstruction are displayed in the right column. The use of navigators prolongs the total scan time, and thus increases the sensitivity to motion, as shown in the single-direction DWI reconstructed with navigated shot phase. The retrospectively self-gated reconstruction discards navigators, and renders sharper DWI. Compared to LLR, ZSSSL is advantageous in resolving clearer tissue boundaries in DWI, as indicated by red arrows.

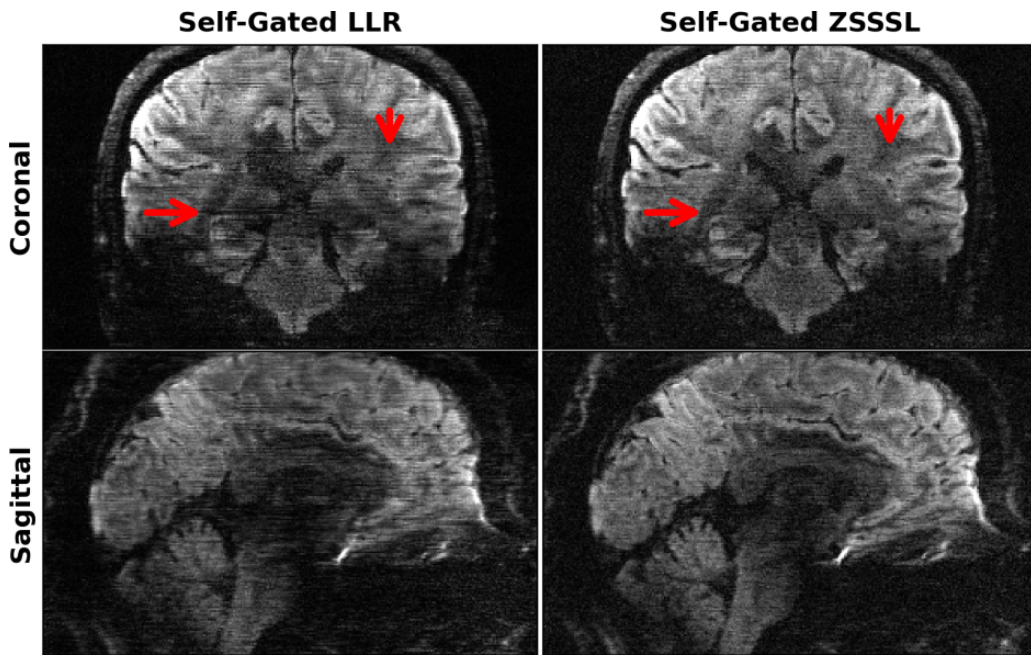


Fig. 3. Single-direction DWI at 0.7 mm isotropic resolution as reconstructed by retrospectively self-gated (left) LLR and (right) ZSSSL in (top) the coronal and (bottom) the sagittal views. The same diffusion direction as in Figure 2 is chosen for display. ZSSSL reduces phase ambiguities in the shot-combined reconstruction, thereby rendering clearer tissue delineation and reduced stripping artifacts (as indicated by the red arrows).

iEPI acquisition and high-resolution DWI reconstruction. The self-gated ZSSSL achieved whole brain diffusion encoding in 21 directions with a b -value of 1000 s/mm^2 at 0.7 mm isotropic resolution, all within a scan time of less than 10 minutes. Technically, this work unrolled ADMM to perform ZSSSL training and testing. Likewise, ADMM was employed to solve the inverse problem in Equation (2) with LLR regularization. This approach assures fair comparison among different regularization methods.

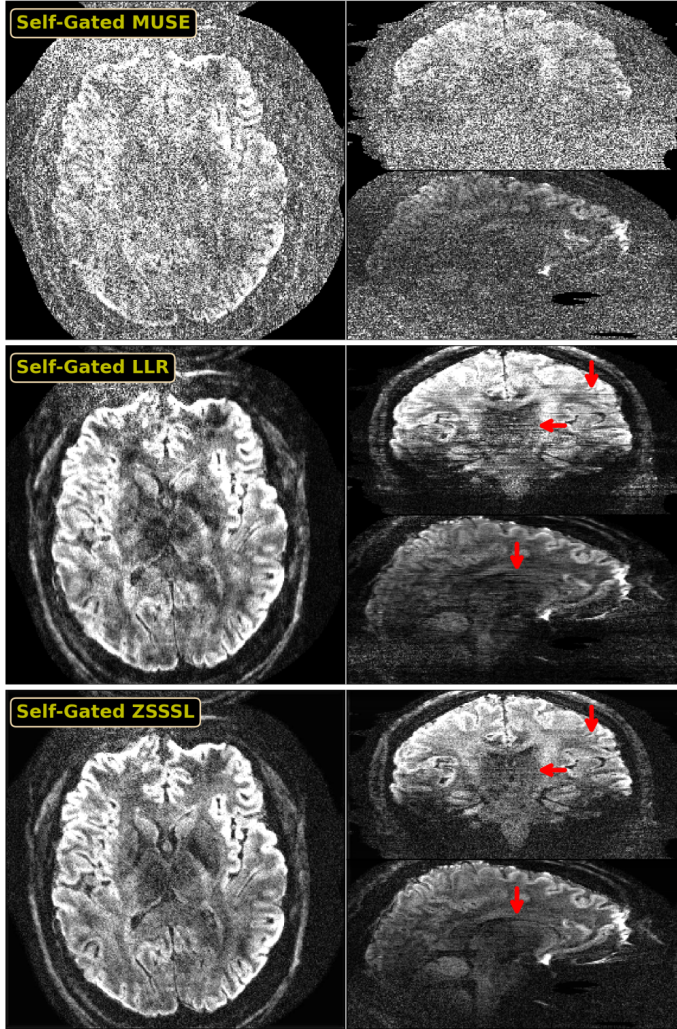
The proposed self-gated ZSSSL approach is well-suited for online reconstruction deployment. Firstly, it requires much shorter acquisition time than the conventional MUSE approach with fully-sampled iEPI and our previous NAViEPI method.

Secondly, ZSSSL does not require large-scale fully-sampled data for training. Instead, the training of ZSSSL is scan specific. Last but not the least, the trained ZSSSL model is applicable to different undersampling factors and to different slices. Fourth, the inference time of ZSSSL is much shorter compared to the LLR regularization approach.

We observed that stripping-type motion artifacts occurred more frequently with sub-millimeter isotropic resolution DWI. In addition, sub-millimeter isotropic voxel resulted in higher noise in DWI. This makes sense, as scans with reduced slice thickness are more susceptible to shot-to-shot phase variations. To enable sub-millimeter mesoscale DWI, Setsompop et al. [45] proposed the gSlider technique with slice phase-dither

0.7 mm mesoscale DWI with 21 volumes @ 10 minutes

(A) Single-dir. DWI



(B) Mean DWI from 20 directions

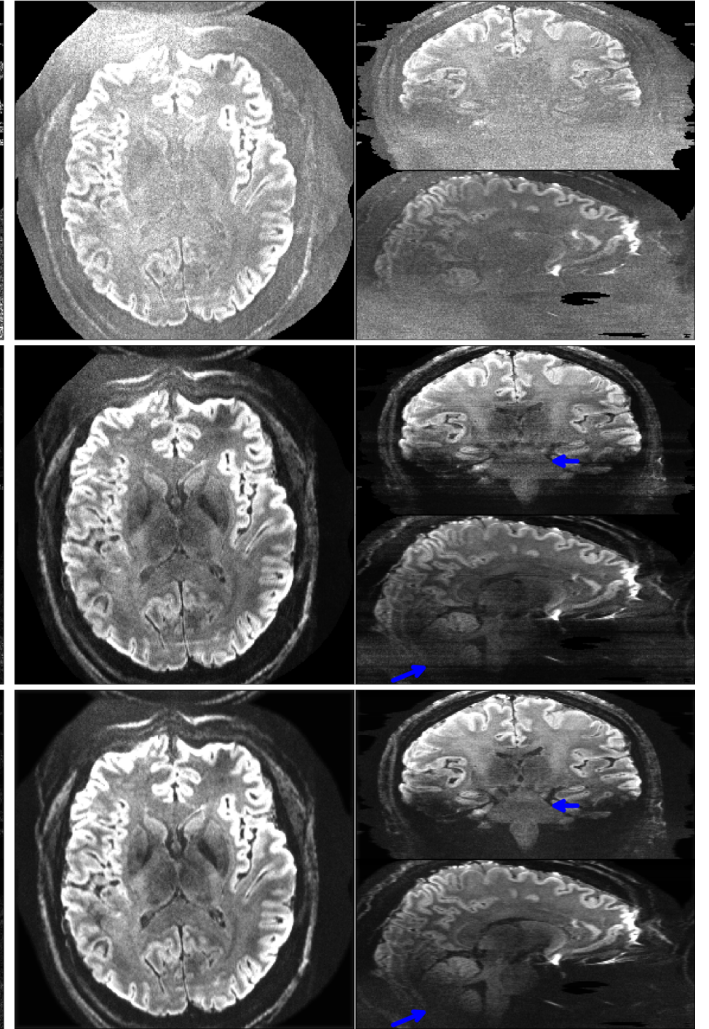


Fig. 4. 0.7 mm isotropic mesoscale DWI with 21 volumes at about 10 minutes without the acquisition of navigators. (A) Single-direction DWI and (B) Mean DWI of 20 diffusion directions at three orthogonal orientations are displayed. DWIs were reconstructed by (top) MUSE, (middle) LLR, and (bottom) ZSSSL. MUSE suffers from severe noise artifacts at such small voxel size. LLR is able to clean up most of the noise, but is still hampered by signal void artifacts in the axial view, which appears as stripping artifacts in coronal and sagittal views (as indicated by red arrows). ZSSSL significantly reduces both noise and signal voids. In the mean DWI images, LLR shows amplified noise in the cerebellum region, whereas ZSSSL yields more homogeneous signal distributions.

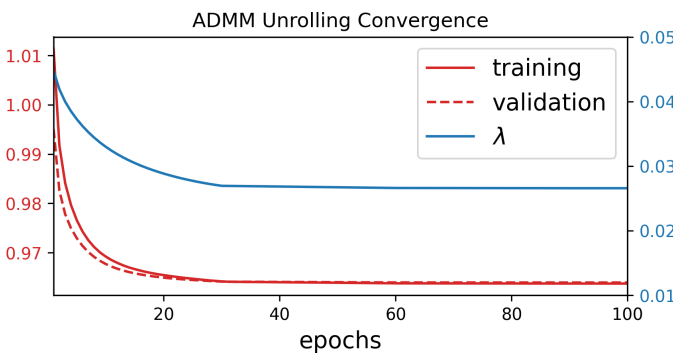


Fig. 5. Convergence analysis along the ADMM unrolling training and validation epochs for the ZSSSL results in Figure 4. Displayed curves are (red solid) the training loss, (red dashed) the validation loss, and (blue solid) the learned regularization strength λ , respectively.

encoding, which excites one slab multiple times with complementary slice encoding schemes. gSlider has been proven effective in alleviating motion sensitivity, because the thicker slab (in comparison to the thin single slice) reduced inter-slice motion. Meanwhile, Hadamard encoding of the slices within a slab gained SNR in the linear inverse reconstruction. However, it has been reported that gSlider has stricter requirements on B_0 and B_1 field homogeneity and shows residual slab boundary blurring [46]. In contrast, the proposed self-gated ZSSSL method requires no such advanced slab encoding, while achieves sub-millimeter resolution at a clinical feasible reconstruction time. Thus, the proposed method can be useful for the probe to high-resolution brain micro-structures in the human connectome project [47].

This work demonstrated the capability of self-gated ZSSSL in reconstructing 0.7 mm isotropic resolution iEPI DWI

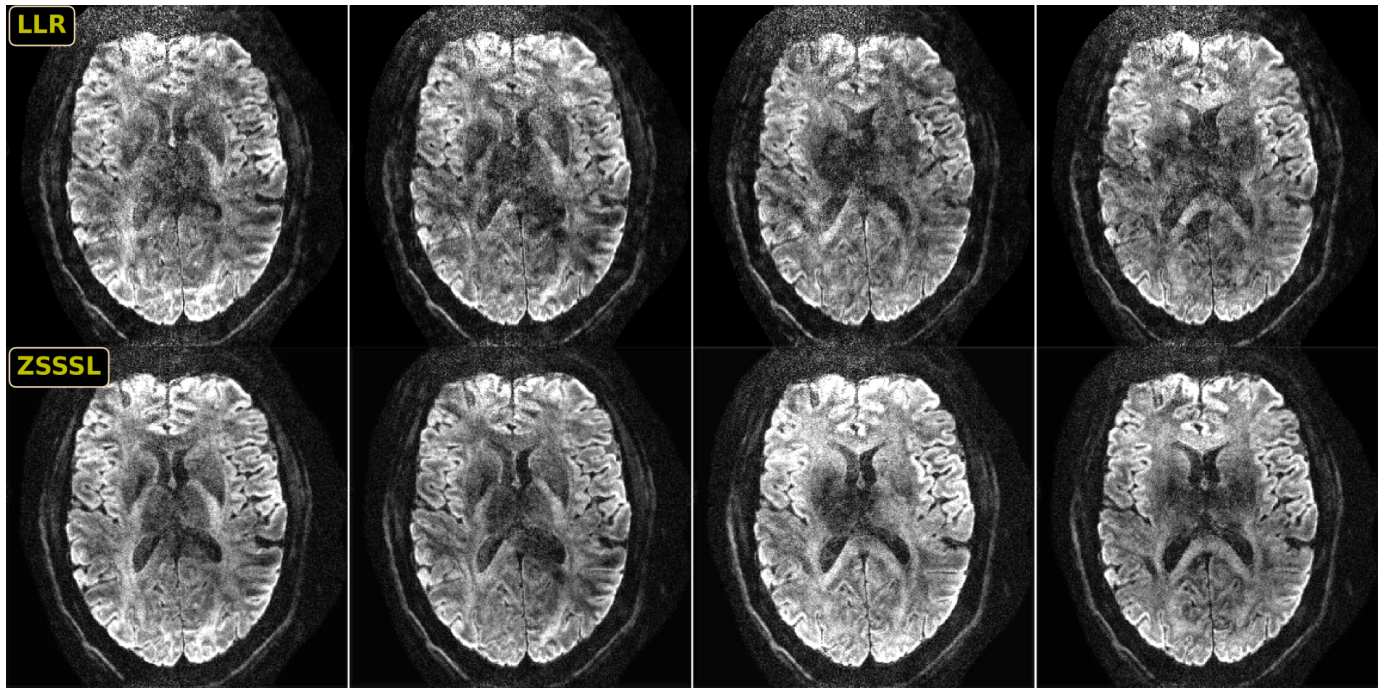


Fig. 6. Prospectively self-gated DWI reconstruction results at 0.7 mm isotropic resolution. Displayed images are one axial slice at four different diffusion-encoding directions. ZSSL enables much cleaner delineations of diffusion contrasts than LLR.

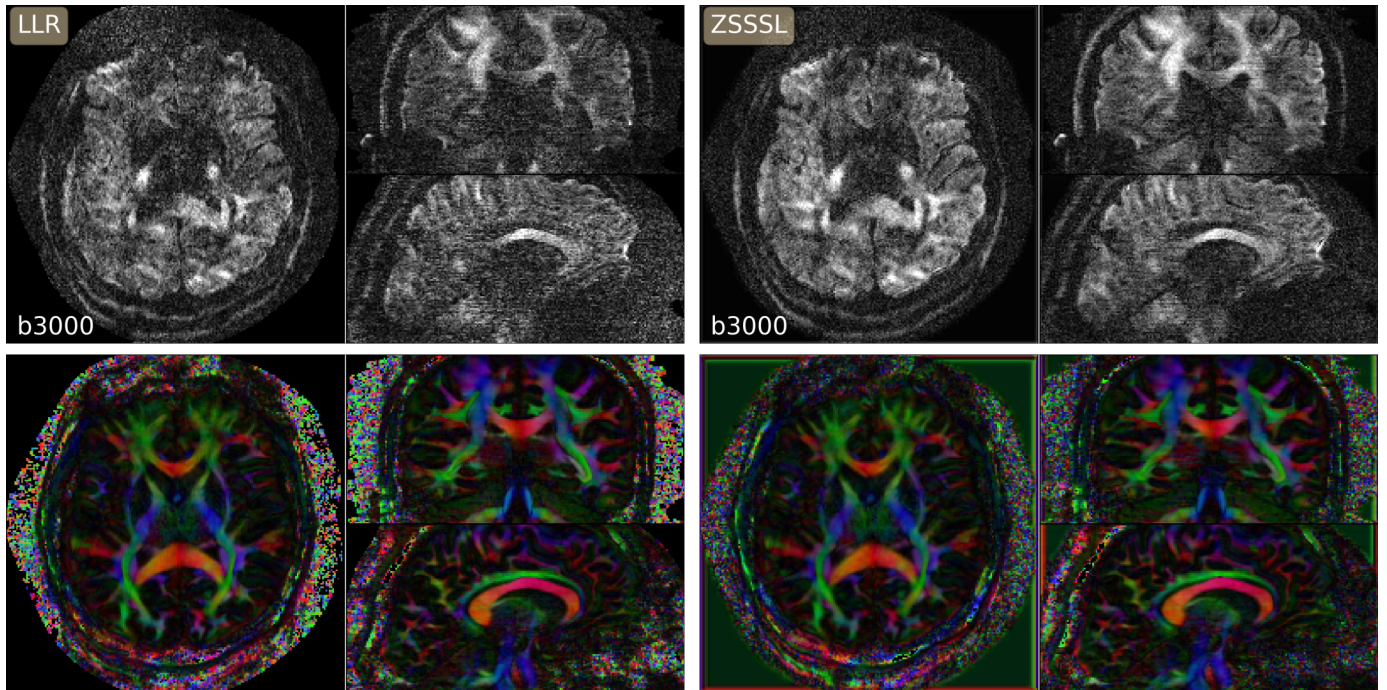


Fig. 7. Reconstruction results of 3-shell 1.0 mm isotropic resolution DWI with 3×3 acceleration: (left) LLR and (right) ZSSL. (Top) Single-direction DWI at the b -values of 3000 s/mm^2 and (bottom) colored fractional anisotropy maps.

with (6×2) -fold acceleration per shot. However, we also observed that the self-gated approach failed to recover aliasing-free DWI in the case of higher acceleration factors (e.g. the $0.5 \times 0.5 \times 2.0 \text{ mm}^3$ DWI data with an acceleration of 10×2 per shot). To address this issue, employing optimized trajectories with a more densely-sampled k -space central region could help better estimate shot phase variations [36], [48].

One limitation of this work is that the 3-shell 126-direction data were trained and inferred in three partitions because such large data cannot be allocated into one single GPU. One potential direction would be to incorporate latent models [49] to reduce the data dimension, which eventually allows efficient representation of large high-dimensional data.

This work did not incorporate off-resonance correction in the reconstruction. As a logic extension, the multi-shot sequence can be modified to encode dynamic B_0 field variation, which can then be employed in the SENSE-based forward operator and reconstruction. An established approach is known as the blip-up/down encoding [50]. This approach can potentially be combined with the model-based reconstruction for joint reconstructions of DWIs and B_0 field maps [51].

VI. CONCLUSION

In this work, we proposed a self-gated zero-shot self-supervised learning reconstruction framework based on ADMM unrolling for high-resolution and motion-robust DWI.

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